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APPENDIX E
SUPPLEMENTAL EVALUATION OF FATE AND TRANSPORT PROCESSES

### E.0 FATE AND TRANSPORT PROCESS

Contaminant fate and transport in the environment may be controlled by a number of factors such as the chemical and physical properties of the contaminants, the geologic formations encountered, the hydrogeologic conditions (vertical and horizontal groundwater gradients, conductivity of the aquifers, and seasonal fluctuations in the groundwater table), precipitation, tidal fluctuations, and topography of the study area. By evaluating the patterns of contaminant distribution and the transport pathways available, it is possible to identify key factors governing contaminant migration within the OU3 study area.

Chemical and biological processes may cause the transformation into other compounds, affecting the fate and transport of contamination. Chemical or biological degradation may be a significant factor for contaminant fate and transport within the OU3 study area; however; the results of the RI concentrations in various environmental media have been fairly constant over time for the contaminants within the OU3 study area. In addition, several of the contaminants are either difficult to biodegrade, or do not transform under normal environmental conditions (i.e. asbestos, PCBs, some metals). Therefore, no further evaluation of chemical or biological processes will be discussed in this RI.

Section E.1 summarizes the dominant transport mechanisms in the environment. Section E.2 discusses the properties of the chemical contaminant which influence their fate and transport. Section E.3 summarizes the contaminant sources and releases to the environment within the OU3 study area associated with historical disposal and operational practices associated with the Raymark Facility. Section E.4 provides a discussion of the physical states of contaminants in the various environmental media within the OU3 study area.

#### E.1 Major Fate and Transport Processes

Contaminants in natural systems are subject to a variety of physical and chemical interactions that influence their fate, and subsequent transport, in the environment. The basic processes governing contaminant transport include advection, diffusion and dispersion, decay, and retardation. Additional mechanisms support fate and transport of chemical contaminants on a smaller scale. The processes are described below.

#### E.1.1 Advection

Advection is the bulk movement of a dissolved-phase or particulate contaminant within a natural fluid flow (air or water) in response to a potential gradient such as pressure (hydraulic head), temperature, or concentration. In the environment, advection is the dominant transport mechanism controlling the physical transport of contaminants. Under advective transport, a contaminant present in air or water is passively carried at the same velocity as the fluid. Examples of advective transport include migration of: contaminant plumes in groundwater downgradient of unlined lagoons, contaminant plumes in surface water downstream of outbreaks to rivers and streams, dust clouds/air emissions downwind on an uncontrolled excavation, and bulk movement of contaminants as the result of tidal fluctuations.

## E.1.2 Diffusion and Dispersion

In environmental systems, diffusion and dispersion are often considered concurrently in the transport of chemical contaminants through plume spreading and reduction in peak concentrations over time. However, the two terms represent distinct transport processes.

Diffusion is generally viewed as the slow migration of dissolved contaminants, in response to a concentration gradient, from areas of higher concentrations to areas of lower concentrations. Because this transport process occurs on the molecular scale, its affect is typically small in comparison to the forces driving the more rapid advective processes. Diffusion, however, may be significant in aquifers that have clayey sands and silts with low groundwater velocities; and it may be the dominant transport process for the movement of soil gas. Diffusion also plays a role in adsorption and other chemical properties discussed in Section 5.2.

Dispersion, on the other hand, is a mechanical process resulting from fluid-flow irregularities which cause contaminants to follow an indirect path from their point of origin. These tortuous and intertwining pathlines (caused by rough spots on or within the ground surface or a pipe wall) lead to mixing and spreading of the contaminants. The flow path irregularities may also arise at faster velocities and under conditions of turbulent flow. In groundwater systems, the contaminant pathway becomes more convoluted as the degree of subsurface heterogeneities

increases. Typical heterogeneities affecting groundwater flow include changes in porosity, hydraulic conductivity, and layering in the aquifer.

Examples of transport by diffusion and dispersion include movement of contaminants within: confined and unconfined aquifers, vadose zone groundwater, surface water, and the atmosphere.

#### E.1.3 Decay

Decay of organic chemical contaminants occurs by biological and non-biological means. In biological decay or biodegradation, chemical transformations are mediated by the extracellular enzymes synthesized by microorganisms residing in soil and aquatic media. These enzymes may be produced for the sole purpose of breaking down organic contaminants as a food source for the microorganisms (metabolism) to catalyze chemical decomposition for some other purpose (cometabolism). Mechanisms of non-biological decay include hydrolysis, photolysis, oxidation, reduction, and methylation. Photolysis, or photo-oxidation, is limited to situations where sunlight is present at the soil surface and in the upper few feet of most surface water.

Although various decay mechanisms may be important in specific environments, biodegradation often comprises a significant portion of the overall decay rate. Chemicals that were not originally present at the site may be produced. In some cases, these daughter products such as vinyl chloride may be more toxic than the original compounds released. Alternatively, biodegradation can significantly reduce the chemical hazards at a site by transforming chemical contaminants to harmless constituents, such as carbon dioxide and water.

Furthermore, depending on the microorganisms and contaminants present, biodegradation can occur under aerobic or anaerobic conditions. However, due to its slow rate in the absence of artificial stimulation, aerobic biodegradation is rarely observed at hazardous waste sites. Anaerobic biodegradation is a much more common occurrence.

#### E.1.4 Retardation

Adsorption-desorption reactions lead to the retardation of contaminant plume migration in groundwater flows, as well as of contaminant movement along the boundaries of surface water flows. Contaminants, moving by advective transport in the direction of the groundwater flow, are slowed down due to the presence of chemical interactions. The cause of this behavior is based on the concept that during some of the contaminant's travel time, a given contaminant may become adsorbed to stationary soil particles and, therefore, will not travel along with the bulk groundwater flow. Only upon desorption will a contaminant rejoin the groundwater flow. As a result, the overall travel time of a groundwater contaminant will increase due to the time spent stationary while adsorbed on soil particles. This retardation phenomenon can also occur in contaminant movement in surface water flows. Surface water contaminants that come into contact with sediment lining the flow path may also become adsorbed. As with groundwater contaminants, adsorption of organic compounds onto sediment particles in the surface water flow is influenced by the organic carbon content of the sediment; adsorption of metals is also influenced by the presence of metal oxide coatings on the soil/sediment particles.

#### E.1.5 Additional Mechanisms

Additional mechanisms, generally secondary in importance, affect fate and transport on a smaller scale. These mechanisms include partitioning, dissolution and precipitation, and biotic uptake.

<u>Partitioning</u> - Chemical contaminants are separated and distributed between various environmental media based on a variety of partitioning coefficients adopted from various engineering disciplines. These include an air-water partitioning coefficient based on the Henry's Law constant (K), an octanol-water partition coefficient (K<sub>ow</sub>) depicting the ratio of solute concentration in octanol (an organic solvent) to that in water, and a soil-water partition coefficient (K<sub>d</sub>) derived from the K<sub>ow</sub> and the amount of carbon-containing material in the soil. These partition coefficients, based on the chemical properties of a particular contaminant, are described in Section 5.2.

#### Dissolution and Precipitation

In the environment, water naturally contains detectable quantities of a number of inorganic constituents (metals and salts). These constituents may be present in precipitation or may be the result of surface water runoff, or groundwater contact with, and dissolution of soil particles and rock surfaces. The amount of dissolved species present depends on the solubility of the individual constituent, as well as on the time and degree of contact between the water and soil/rock surfaces. Furthermore, once in solution, dissolved constituents may return to their solid form by precipitation onto soil/rock surfaces.

Under equilibrium conditions, the balance between precipitation and dissolution of an inorganic constituent in a natural water system depends on a number of factors, including:

Acid/Base Chemistry - The balance between H<sup>+</sup> and OH<sup>-</sup>, expressed as pH, is a dominant control on aqueous solubility of many inorganics. Typically, the aqueous solubility of many metals increases as pH decreases and the water becomes more acidic. These effects are most notable in areas surrounding mine tailing dumps, as well as in areas where large quantities of acidic wastewater have been disposed.

Oxidation State - The presence of dissolved oxygen and other oxidants influences the ratio of stable states of various inorganics and, therefore, impacts their overall solubility. Reduction-oxidation or redox potentials can also be influenced by biological activity related to contaminant degradation. For example, the dissolution of iron in the environment is strongly influenced by redox reactions which are both biologically and non-biologically mediated.

<u>Bulk Chemistry</u> - The concentration of dissolved inorganics and other solutes determines the type of chemical compounds and complexes that can form, and potentially affects their overall stability. In addition, the total concentration of dissolved ions or ionic strength can affect the solubility of chemical contaminants.

<u>Complexation</u> - Certain soluble organics can form strong chemical bonds to metal ions in solution forming chelates and coordination complexes. These complexes tend to "tie

up" the metal species either in solution thereby preventing them from being adsorbed or precipitated onto soil/rock surfaces, or in an insoluble form that precipitates readily.

## Biotic Uptake

A limited quantity of chemical contaminant fate and transport may also be the result of biotic uptake in plant and animal tissue. Typically, organic compounds are absorbed and retained by biotic materials depending on the individual compound's solubility in lipids (fats and oils) present in the plant and animal tissue. Because many of these lipid-soluble compounds have poor solubilities in water, the organic compounds that are easily adsorbed onto organic carbon in soil are also easily accumulated in plant tissue. Uptake of metals, on the other hand, generally increases with increased water solubility and decreased ability to adsorb onto soil particles.

Biotic uptake in plants primarily occurs through the root system. However, adsorption of aerially-deposited dusts and aerosols through plant leaves may also be significant. For most animals, contaminant uptake occurs through the food chain. However, benthic organisms (those living in the bottom sediments of a water body) appear to take up contaminants through direct ingestion and physical contact.

## E.2 Chemical Properties that Influence Contaminant Fate and Transport

Several chemical properties strongly influence the fate and transport of contaminants in the environment. Each property, along with its relationship to the various fate and transport properties, is defined below.

#### E.2.1 Water Solubility

Water solubility is one of the primary chemical characteristics used to assess chemical fate and transport in the environment. The water solubility of a chemical contaminant provides considerable insight into its mobility, stability, tendency to adsorb to soil or sediment, and propensity to accumulate in the environment or bioaccumulate in the food chain.

Chemicals with high water solubilities, greater than 1000 mg/l or 0.1 percent (Ney, 1990), will have a tendency to remain dissolved in the water column and will not likely partition to soil/sediment particles or accumulate in the environment or bioaccumulate in aquatic organisms in the food chain. Highly soluble chemicals are less likely to volatilize from water and are generally more prone to biodegradation and metabolism. As these characteristics indicate, chemicals with higher water solubilities are more likely to be mobile, and therefore less likely to persist, in the environment.

Conversely, chemicals with low water solubilities, less than 10 mg/l or 0.001 percent (Ney, 1990), are generally less mobile and are more likely to adsorb to soil/sediment particles, accumulate and/or bioaccumulate, and persist in the environment. Low solubility chemicals tend to biodegrade or be metabolized less readily in plants and animals.

Many of the VOCs identified during the RI have moderate to high water solubilities. These included: the ketones with the highest solubilities (ranging from very high to miscible), most of the chlorinated compounds with high solubilities, and the aromatic compounds generally with moderate solubilities. The ketones would, therefore, more readily enter into an aqueous solution and migrate in groundwater and surface water than other VOCs. The remaining less soluble VOCs, the SVOCs, with pesticides/PCBs, the dioxins/furans, the metals, and asbestos detected in the study area are more likely to stay adsorbed to soil and sediment particles and be less mobile.

#### E.2.2 Octanol-Water Partition Coefficient (Kow)

The octanol-water partition coefficient (K<sub>ow</sub>) is a measure of the tendency of an organic compound to partition between organic (in this case, octanol) and aqueous phases. The chemical-specific K<sub>ow</sub> is an indicator of its water solubility, mobility, sorption, and bioaccumulation. It has also been shown to correlate well with bioconcentration factors in aquatic organisms and the adsorption to soil and sediment (Howard, 1990).

The higher the K<sub>OW</sub> value of a specific chemical, the greater is the chemical's potential for sorption to soil/sediment particles, the lower its mobility, and the more likely it is to accumulate

in the environment and bioaccumulate in the food chain. A high  $K_{ow}$ , greater than 1000 (Ney, 1990), is indicative of low water solubility and greater persistence in the environment.

Conversely, the lower the  $K_{OW}$  value, the greater is the chemical's potential to biodegrade and be metabolized by plants and animals. A low  $K_{OW}$ , less than 500 (Ney, 1990), is indicative of high water solubility, high mobility, and little or no accumulation or bioaccumulation.

The VOCs identified in the study area have widely varied  $K_{OW}$  values. These included: the ketones with the lowest  $K_{OW}$  values; the chlorinated compounds with generally low  $K_{OW}$  values, with the exception of tetrachloroethene (PCE) with a high  $K_{OW}$ ; and the aromatic compounds generally with low to moderate  $K_{OW}$  values, except for ethylbenzene with a high  $K_{OW}$ . The VOCs, as a group, have a stronger tendency to enter into an aqueous solution and migrate in groundwater and surface water than the other organic compounds. Most other organic compounds detected in the study area have high  $K_{OW}$  values, indicating a stronger preference to adsorb to soil/sediment particles and be less mobile. Metals and asbestos also preferentially adsorb to soil/sediment particles, due more to their low water solubilities rather than their ability to partitioning between organic and aqueous phases.

#### E.2.3 Adsorption Partition Coefficient (K<sub>d</sub> & K<sub>oc</sub>)

The adsorption partition coefficients ( $K_d$  and  $K_{OC}$ ) are measures of the tendency of an organic contaminant to bind to soil or sediment particles. Adsorption coefficients are useful in evaluating the mobility of contaminants in the subsurface media.  $K_d$  is an experimentally derived coefficient which measures the tendency of a chemical to sorb to a particular soil/sediment media. The  $K_d$  is both a chemical- and a sorption media-specific coefficient related to the organic carbon content of the media.  $K_{OC}$ , the organic carbon partition coefficient, is a chemical-specific value derived by dividing  $K_d$  by the organic carbon content of the sorption media in order to normalize the value to the organic carbon content of the media.  $K_{OC}$  is defined as the ratio of the amount of chemical absorbed per unit weight of organic carbon in the media to the concentration of the chemical in solution at equilibrium.

Chemicals with high adsorption coefficients have a tendency to bind to soil/sediment particles containing organic carbon and are, therefore, relatively immobile in groundwater. A relationship developed between an organic compound's K<sub>oc</sub> and its mobility is as follows (Dragun, 1988):

K <sub>oc</sub> (ml/g)	Mobility Class
>2,000	Immobile
500-2,000	Low Mobility
150-500	Intermediate Mobility
50-150	Mobile
<50	Very Mobile

Most of the VOCs identified in the study area have low to moderate  $K_{OC}$  values. These included: the ketones with the lowest  $K_{OC}$  values; and the chlorinated and aromatic compounds with low to moderate  $K_{OC}$  values, with the exception of PCE with a high  $K_{OC}$ . The VOCs, as a group, have a low tendency to adsorb to soil/sediment particles and are, therefore, more mobile in the environment than the other organic compounds. Most other organic compounds detected in the study area have high  $K_{OC}$  values, indicating a tendency to sorb to soil/sediment particles and be more persistent in the study area. Metals and asbestos also preferentially adsorb to soil/sediment particles, due in large part to their low water solubilities.

#### E.2.4 Vapor Pressure

Vapor pressure is a measure of the ability of a compound volatilizing from the pure liquid phase into the vapor phase. It provides an indication of the rate at which a chemical will volatilize from soil, sediment, or water. This property is of primary significance at environmental interfaces, such as surficial soil/air, surficial sediment/air, and surface water/air.

Chemicals with higher vapor pressures are expected to enter into the vapor phase more readily than those with lower vapor pressures. If chemicals with low vapor pressures (less than  $10^{-6}$  mm Hg) are present in the atmosphere, they are most likely associated with adherence to suspended particulate matter.

VOCs all have relatively high vapor pressures and will, therefore, tend to enter the vapor phase in the presence of air. Vapor pressures for the major groups of VOCs identified in the study area (ketones, chlorinated hydrocarbons, and aromatic hydrocarbons) are generally many orders of magnitude higher than vapor pressures for SVOCs, pesticides/PCBs, and dioxins/furans. The VOCs identified in the RI are likely to volatilize from the soil into the soil pore space in the vadose zone, and from surface water into the ambient air. Metals and asbestos, having very low or immeasurable vapor pressures, do not readily volatilize.

## E.2.5 Henry's Law Constant (K)

Henry's Law Constant (K) is an air-water partition coefficient which serves to indicate the tendency of a chemical to volatilize or evaporate from water. K can be expressed as dimensionless (relating the concentration of a chemical in the vapor phase to its concentration in water at equilibrium conditions) or with units (representing the ratio of the vapor pressure of the chemical in atmospheres to its water solubility in moles/m³).

In general, chemicals with low K values (less than 5 x 10<sup>-6</sup> atm·m³/mole) have a low tendency to volatilize from surface water into the ambient air or from groundwater into the soil pore spaces. For chemicals with higher K values, volatilization and diffusion into the ambient air and/or soil pore spaces could be a significant fate and transport mechanism (Dragun, 1988).

The VOCs identified in the study area all have moderate to high K values. These included: the ketones with moderate K values; and the chlorinated and aromatic compounds with high K values. The VOCs would, therefore, be likely to volatilize from the aqueous phase to the vapor phase. Most SVOCs, pesticides/PCBs, and dioxins/furans detected in the study area have low to moderate K values, indicating they are less likely to volatilize from the aqueous phase. Metals and asbestos, having very low or immeasurable K values, do not readily volatilize from the aqueous environment.

#### E2.6 Specific Gravity

Specific gravity is defined as the ratio of the weight of a given volume of a pure chemical at a specified temperature to the weight of the same volume of water at a given temperature.

Specific gravity indicates the tendency of a chemical to float or sink in water given that it is present in its pure form or at a very high concentration.

For contaminants that exist in a separate phase from the water, those with specific gravities less than 1 (such as aromatic hydrocarbons and ketones) will tend to rise or float to the top of the water column as a sheen or top layer. Contaminants with specific gravities greater than 1 (such as most chlorinated hydrocarbons, pesticides/PCBs, and dioxins/furans) tend to mix and/or sink in the water column. Since metals and asbestos generally have low water solubilities, and do not readily exist in the environment as a separate phase, they due not tend to either rise or sink in the water column based on their specific gravities. Instead, they may rise or sink along with particles to which they might adsorb.

#### E.3 Physical State of Contaminants in Environmental Media

The physical state in which contaminants exist in various environmental media dictate the manner in which they may migrate or be transported. The phases in which organic and inorganic contaminants may be present in the sediment/soil and surface water/pore water within the OU3 study area, as well as the potential to migrate from these phases, are discussed below. Issues related to groundwater contaminants are addressed in a qualitative fashion as it relates to the migration of contaminants to or from soil/sediment and surface water.

#### E.3.1 Soil/Sediment Contaminants

Organic and inorganic contaminants introduced at the ground surface through spills, discharges, or disposal may contaminate the underlying soil/sediment, migrate to downgradient groundwater and surface water/pore water sources, and/or be discharged into the atmosphere. These contaminants may be present in soils and sediment in three principal physical phases: solid, liquid, or vapor.

<u>Solid Phase</u> - Inorganic contaminants (metals and asbestos) are often present in the solid phase, as bulk solids or as suspended particulates in aqueous discharge, when disposed on the surface or into the subsurface. In the solid phase, inorganics are relatively immobile in a

soil or sediment matrix unless the particle sizes are sufficiently small that migration may occur as a suspended particle or colloid. Solid phase metal contaminants may also be broken down and/or leached by precipitation or subsequent liquid spills that solubilize the inorganic constituents, and may thereby become mobile.

Organic contaminants are rarely present in the environment in the solid phase.

Liquid Phase - Most of the organic contaminants related to industrial sites are present in the liquid phase (as a separate product or a mixture of liquid chemicals) or in the aqueous phase (dissolved in a water-based solution). Once these liquids enter unsaturated soil or sediment, several processes may occur. The organic contaminants may: (1) become adsorbed to the soil/sediment matrix because of low water solubilities, high soil/sediment organic carbon partition coefficients, and/or high soil/sediment organic content; (2) remain in a liquid phase in the soil/sediment pore water spaces as a result of pore size and interfacial tension; or (3) volatilize into soil/sediment pore spaces if the contaminants have high vapor pressures or high air-liquid partition coefficients. If present in sufficient quantity, a liquid contaminant may migrate downward under the influence of gravity and enter underlying groundwater.

Metals may also be present in the liquid phase, as dissolved ions. In low pH conditions, more metals are apt to be present as dissolved ions and are, therefore, more readily available to mobilize. For instance, aqueous acid solutions used for cleaning metal surfaces contain numerous dissolved metals. If the solution is discharged into soil/sediment, some of the metals may remain in the aqueous phase and/or additional metals present in the soil/sediment may dissolve, if the pH is sufficiently low. In addition, some metals may precipitate from solution in the presence of minerals in the soil/sediment matrix and become adsorbed onto soil/sediment particles.

Asbestos is not present in the environment in the liquid phase.

<u>Vapor Phase</u> - Organic contaminants, principally VOCs, present in an unsaturated soil or sediment matrix may volatilize into the pore spaces and migrate through the soil/sediment if the vapor pressure is sufficiently high. Once in the vapor phase, VOCs can migrate readily if no barriers impede their movement.

Other organic contaminants (with lower vapor pressures), as well as inorganic contaminants, are not typically present in the environment in the vapor phase.

#### E.3.2 Surface Water Contaminants

Organic and inorganic contaminants introduced at the ground surface through spills, discharges, or disposal may contaminate the surface water, migrate to underlying soil/sediment and groundwater sources, and/or be discharged into the atmosphere. These contaminants may be present in surface water in three principal physical phases: solid, liquid (aqueous or non-aqueous), or vapor.

<u>Solid Phase</u> - Metals, asbestos, and some organic compounds that are either insoluble or otherwise remain undissolved in water may migrate in the surface water as suspended particulates or colloids. Metal ions may also adsorb to the surface of colloidal particles, through surface attractions, and migrate through the surface water.

Aqueous Liquid Phase - Dissolved aqueous phase organic and inorganic contaminants present in surface water may be transported simultaneously by three processes: advection, mechanical dispersion, and molecular diffusion. During transport, contaminants may interact with other aqueous-phase contaminants and ions. A variety of chemical, physical, and biological interactions can significantly retard or accelerate the rate of contaminant transport, or transform the contaminants into other chemicals or phases.

Undissolved organic and inorganic contaminants, as well as asbestos, are not present in the environment in the aqueous liquid phase. In surface water or groundwater aquifers, they are generally present as solid particulates or colloids.

Non-Aqueous Liquid Phase - Non-aqueous phase liquids (NAPLs), such as organic solvents or fuels containing little or no water, may exist in surface water as a separate/immiscible liquid. Contaminants present in this phase behave differently than those dissolved in the aqueous phase. NAPLs are a continuing source of dissolved-phase organics to surface water and groundwater. Because NAPLs have limited surface areas relative to their volume, dissolution may extend over an indefinite amount of time.

Dense NAPLs (DNAPLs), such as TCE which has a higher density than water, tend to migrate vertically downward under the influence of gravity in response to both the physical properties of the contaminant and the characteristics of the surface water (or underlying groundwater aquifer). Contaminant properties influencing transport include density, viscosity, and interfacial tension. Groundwater aquifer characteristics influencing transport include pore size, permeability, general stratigraphy (location and topography of low-permeability units), and micro-structures (such as small fractures in subsurface soil media or bedrock) (EPA, 1992).

Light NAPLs (LNAPLs), such as toluene which has a lower density than water, are generally present near the surface of surface water sources (or at the water table surface of groundwater aquifers). LNAPLs tend to migrate laterally along the top of the water surface in response to the organic properties and groundwater aquifer characteristics described above for DNAPLs.

Metals and asbestos are not typically present in the environment in the non-aqueous liquid phase. If present in surface water or groundwater aquifers, they are generally found as solid particulates or colloids.

<u>Vapor Phase</u> - Organic contaminants, principally VOCs, present in surface water may volatilize into the atmosphere if the vapor pressure is sufficiently high. Once in the vapor phase, VOCs can migrate readily if no barriers impede their movement.

Other organic contaminants with lower vapor pressure, as well as inorganic contaminants, are not typically present in the environment in the vapor phase.

# E.4.1 Contaminant Fate and Transport in Sediment and Surface Water – Ferry Creek and Adjacent Wetlands

Various chemical contaminants are present in the sediment and surface water in the Ferry Creek channel, and in sediment in the wetlands adjacent to the channel, the western reaches of the ditch extending from Ferry Creek westward between properties located at 250 and 230 Ferry Boulevard, and in the single location sampled in the triangular-shaped property between East Broadway and Ferry Boulevard. Several pathways are available for contaminants to have

entered these media, and for them to subsequently be transported downstream and/or to adjacent media.

The chemical and physical properties of the chemical contaminants affects their ability to migrate into the sediment and surface water in the Ferry Creek Area, and to subsequently be transported to adjacent media. Contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend adsorb to the sediment particles. These contaminants are apt to remain localized in undisturbed sediment, or to migrate on sediment particulates transported by surface water flows, rainwater runoff, tidal fluctuations, or groundwater discharge. Contaminants adsorbed to sediment particles may later become desorbed and/or dissolved following contact with solvents or acids, if present. Contaminants with low soil/sediment adsorption coefficients, high water partition coefficients, or high water solubilities would tend to remain in solution and be free to migrate with naturally occurring waters.

Groundwater discharge to the creek, tidal fluctuations, and seep areas along the Ferry Creek channel may also cause contaminants to be transported into and/or pass through the sediment. Again, those contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend to adsorb to the sediment particles; while those with low soil/sediment adsorption coefficients, high water partition coefficients, and high water solubilities would tend to remain in solution and be free to migrate with natural water sources. Groundwater discharges and seeps containing organic solvents or acids may also cause leaching of chemical contaminants from the surface of sediment particles, and subsequent migration into naturally occurring waters.

The fate and transport of contamination is, therefore, dependent on the type of chemical contaminant present. A discussion of the fate and transport of the various categories of chemical contaminants in sediment and surface water throughout the Ferry Creek Area is presented below. Since no elevated levels of SVOCs or PCBs were present in surface water samples throughout Ferry Creek, and since no samples were collected for analysis of dioxins/furans and asbestos (since these contaminants are virtually insoluble in water), the fate and transport discussion related to surface water is limited to VOCs and metals.

<u>VOCs</u> - Due to their high water solubilities, low soil/sediment partition coefficients, and low octanol-water partition coefficients, VOCs in a sediment/surface water environment tend to remain dissolved in, and free to be transported by, the aqueous media; they do not readily adsorb to sediment particles. Furthermore, VOCs present in surface sediment located in relatively dry locations have more of a tendency to volatilize into the atmosphere than other organic contaminants due to their high vapor pressures. VOCs present in surface water (especially in regions of high turbulence or aeration) also have a tendency to volatilize into the atmosphere due to their high vapor pressures and high air-water partition coefficients.

These chemical properties support the infrequent detection of elevated levels of VOCs in both surface and subsurface sediment throughout the Ferry Creek channel and the adjacent wetland areas, and the corresponding frequent detection of elevated levels of VOCs, primarily chlorinated hydrocarbons, in surface water throughout the length of Ferry Creek.

In the sediment, the highest concentration of an individual VOCs (3,400 ug/kg 1,1-DCA; detected more than an order of magnitude greater than any other VOC) was detected in surface sediment in the Ferry Creek channel just upgradient of the culvert under East Broadway in Area A-1. At this location, the surface water flow was slow moving and the channel was deep, factors may have resulted in deposition of sediment, higher organic-content silts, and/or DNAPLs that could encourage adsorption of VOCs onto sediment/silt particles.

In the surface water, the higher concentrations included: 1,1-DCE, 1,2-DCE, and 1,1,1-TCA in the northern reaches of Ferry Creek near the first major bend in the channel and in the channel extending into the western arm of the wetlands both adjacent to the Morgan Francis property in Area A-1; and 1,1,1-TCA south of the culvert under East Broadway and in the vicinity of an observed seep in the south-central region of Area A-3. These results may have been caused by migration from a nearby source area due to surface runoff or groundwater discharge.

<u>SVOCs (primarily PAHs)</u> - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, SVOCs in a sediment/surface water environment are more readily adsorbed to sediment particles than are VOC contaminants. Furthermore, SVOCs present in surface sediment located in relatively dry locations have a

tendency to volatilize into the atmosphere to some degree. Their ability to volatilize, however, is less than that of VOCs (with higher vapor pressures) but greater than that of PCBs and dioxins/furans (with much lower vapor pressures).

These chemical properties support the frequent detection of elevated levels of SVOCs, primarily PAHs, in surface sediment throughout the Ferry Creek channel and the adjacent wetland areas. The higher frequency of PAH detection in surficial samples, especially in higher deposition areas in the Ferry Creek channel, suggests that the PAHs were adsorbed onto sediment particles that may have migrated to these areas with surface water flows, rainwater runoff, or tidal fluctuations. Less frequent detections in subsurface sediment suggest that the contaminant source is relatively recent (resulting in surficial deposition of contamination) and/or the particles to which the PAHs adsorb have relatively low densities (resulting in frequent resuspension and surficial redeposition).

The highest concentrations of PAHs were detected in surface sediment in the northern reaches of Ferry Creek near the first major bend in the channel adjacent to the Morgan Francis property in Area A-1, and south of the culvert under East Broadway and in the vicinity of an observed seep in south-central region of Area A-3. These results may have been caused by migration from a nearby source due to surface water runoff or groundwater discharge, and subsequent adsorption to sediment particles.

<u>PCBs</u> - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, PCBs in a sediment/surface water environment are more readily adsorbed to sediment particles than are either VOC or SVOC contaminants. Furthermore, PCBs present in surface sediment located in relatively dry locations have little or no tendency to volatilize into the atmosphere due to their low vapor pressures, compared to the highly volatile VOCs and somewhat volatile SVOCs.

These chemical properties support the frequent detection of elevated levels of PCBs, particularly Aroclor 1262 and Aroclor 1268 (with less frequent detection of Aroclor 1248, Aroclor 1254, and Aroclor 1260), in both surface and subsurface sediment throughout the Ferry Creek channel and the adjacent wetland areas. The high frequency of PCB detection in the sediment suggests that the PCBs were adsorbed onto sediment particles that were placed

there as fill material from the Raymark Facility or may have migrated to these areas with surface water flows, rainwater runoff, or tidal fluctuations. The relatively similar frequency of detection in both surface and subsurface sediment in the Ferry Creek channel suggests that deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth) and/or the particles to which the PCBs adsorb have relatively high densities (resulting in infrequent resuspension and redeposition of previously deposited sediment).

The levels of PCBs were relatively consistent in sediment throughout the Ferry Creek channel and adjacent wetlands, with the highest levels detected in both surface and subsurface samples immediately upgradient of the culvert under East Broadway in Area A-1 and immediately downgradient of the culvert in Area A-3.

<u>Dioxins and Furans (as TEF)</u> - As with PCBs, dioxins and furans have low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, resulting in a tendency for the contaminants to adsorb to sediment particles in a sediment/surface water environment. Furthermore, dioxins and furans present in surface sediment located in relatively dry locations have little or no tendency to volatilize into the atmosphere due to their low vapor pressures, again similar to PCBs.

These chemical properties support the frequent detection of elevated TEF levels in surface sediment throughout the Ferry Creek channel, and the somewhat less frequent detection in subsurface sediment from the northern and southern reaches of the Ferry Creek channel. The high frequency of detection suggests that the dioxins and/or furans were absorbed onto sediment particles that migrated to these areas with surface water flows, rainwater runoff, or tidal fluctuations. The relatively similar frequency of detections in surface and subsurface sediment suggests that deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth) and/or the particles to which the dioxins and/or furans adsorb have relatively high densities (resulting in infrequent resuspension and redeposition of previously deposited sediment).

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The highest TEF levels were detected in both surface and subsurface sediment samples immediately upgradient of the culvert under East Broadway in Area A-1, and immediately downgradient of the culvert in Area A-3.

Metals (primarily Copper and Lead) - Metals tend to be less mobile in the environment than organic contaminants. Under natural conditions, dissolution of metals may occur in the presence of water (due to surface water flows, rainwater runoff, or tidal fluctuations) and some metal ions may migrate. However, many of these dissolved metals can readily precipitate out of solution and adsorb to sediment particles. Copper and lead will be more soluble and remain in an aqueous state if the pH is sufficiently low.

These properties support the frequent detection of elevated levels of copper and lead in both surface and subsurface sediment throughout the Ferry Creek channel and adjacent wetlands, and their corresponding infrequent detection in surface water.

In the sediment, the similar frequency of detections in surface and subsurface samples suggests that deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth) and/or the metals migrated in the dissolved state and precipitated on sediment particles upon contact. The highest levels detected in both surface and subsurface samples immediately upgradient of the culvert under East Broadway in Area A-1, and immediately downgradient of the culvert in Area A-3. Elevated concentrations of barium were detected at similar frequencies, with the highest concentrations at the same locations, as copper and lead.

The fact that copper and lead are not apt to dissolve in an aqueous environment under natural conditions (neutral pHs) is supported by their low frequency of detection in surface water from Ferry Creek. In particular, copper and lead were not detected above background concentrations in surface water from north of the culvert under East Broadway, and elevated levels were detected infrequently in surface water from south of the culvert.

Elevated concentrations of barium (being much more water soluble) were detected in surface water throughout the entire length of Ferry Creek.

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Raymark - Ferry Creek, OU3

The highest concentration of these three metals were detected in the high deposition area at the confluence of the Ferry Creek channel with the ditch extended westward between 250 and 230 Ferry Boulevard in Area A-3. These metals may have been present in suspended particulate in this area, or they may have been leached out of contaminated sediment or soils.

It is important to note that elevated levels of lead and/or copper were detected in sediment and soil samples submitted for SPLP analysis from regions north of the culvert under East Broadway (including: five soil samples throughout Area A-1) and south of the culvert (including: three soil samples from the commercial/industrial properties in the northern region of Area A-2, one sediment sample from the northeastern region of the wetland area in Area A-3, and two soil samples in the region north of the Ferry Creek channel in Area A-3). This suggests that these materials may result in a continual release of copper and lead into the surrounding media.

<u>Asbestos</u> - Asbestos is relatively immobile in the environment, except through bulk advective movement or dispersion in aqueous or airborne environments.

Elevated levels of asbestos were detected frequently in both surface and subsurface sediment in the Ferry Creek channel north of the culvert under East Broadway and in the limit number of subsurface sediment samples collected in the triangular-shaped property located between East Broadway and Ferry Boulevard in Area A-1. Less frequent detections were in both surface and subsurface sediment throughout that section of the Ferry Creek channel and adjacent wetlands south of the culvert in Area A-3. The similar frequency of detection in both surface and subsurface sediment in the northern section of the Ferry Creek channel suggests that deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth). The findings further suggest that asbestos may have been transported in discharge from Lagoon No. 4, as well as by transport in rainwater runoff from asbestos-laden fill materials placed in close proximity to the northern section of the Ferry Creek channel.

The highest concentrations of asbestos were detected in surface and subsurface samples north of the culvert under East Broadway along the central portion of the Ferry Creek channel in Area A-1, in subsurface samples from the triangular-shaped property between East

Broadway and Ferry Boulevard in Area A-1, and in surface samples south of the culvert from the Ferry Creek channel and adjacent wetlands in the northern and north-central regions of Area A-3.

# E.4.2 Contaminant Fate and Transport in Soil – Ferry Creek and Adjacent Wetlands

Various chemical contaminants are also present in the soil located at commercial and industrial properties adjacent to Ferry Creek and the wetland areas. Several pathways are available for contaminants to have entered the soil in these regions, and for them to subsequently be transported to adjacent media.

As with the sediment and surface water, the chemical and physical properties of the chemical contaminants in soil affects their ability to migrate in the Ferry Creek Area, and to subsequently be transported to adjacent media. Contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend to adsorb to the soil particles or remain in the soil pore spaces. These contaminants are apt to remain localized in undisturbed soil. Contaminants in soil pore spaces or adsorbed to soil particles may later become mobilized through contact with solvents or acids, precipitation, or groundwater. Contaminants with low soil adsorption coefficients, high water partition coefficients, and high water solubilities would tend to remain in solution and be free to migrate with naturally occurring waters either downward to the water table or laterally to a surface water source.

The fate and transport of contamination is, therefore, dependent on the type of chemical contaminant present. A discussion of the fate and transport of the various categories of chemical contaminants in soil throughout the Ferry Creek Area is presented below.

<u>VOCs</u> - Due to their high water solubilities, low soil partition coefficients, and low octanol-water partition coefficients, VOCs are readily leached from soil and transported into an aqueous media. Furthermore, VOCs present in surface soil have more of a tendency to volatilize into the atmosphere, and VOCs absorbed to soil particles or present in soil pore spaces are more apt to volatilize into unsaturated soil pore spaces or voids and may potentially migrate to the

surface or follow some other preferential flow paths (subsurface drains, utility conduits, etc.), than other organic compounds due to their high vapor pressures.

These chemical properties support the infrequent detection of elevated levels of VOCs in surface soil and the somewhat higher frequency in subsurface soil throughout the Ferry Creek Area. The higher concentrations of VOCs, primarily aromatic hydrocarbons (benzene, toluene, ethylbenzene, and/or xylenes (total)), were detected in subsurface soils located in a few of commercial and industrial properties between Ferry Boulevard and the Ferry Creek channel in Area A-2. The presence of these contaminants may have been the result of localized releases at the properties.

<u>SVOCs (primarily PAHs)</u> - Due to their low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, SVOCs are more readily adsorbed to soil particles than are VOC contaminants. Furthermore, SVOCs present in surface soil have a tendency to volatilize into the atmosphere to some degree, and SVOCs adsorbed to soil particles or present in soil pore spaces have a tendency to volatilize into unsaturated soil pore spaces or voids and may potentially migrate to the surface or follow some other preferential flow paths to some degree. Their ability to volatilize, however, is less than that of VOCs (with higher vapor pressures) but greater than that of PCBs and dioxins/furans (with much lower vapor pressures).

These chemical properties support the frequent detection of elevated levels of SVOCs, primarily PAHs, in surface soil and the somewhat lower frequency in subsurface soil throughout the Ferry Creek Area. The high frequency of detections suggests that the PAHs were sorbed onto soil particles placed in the area as fill material. The somewhat higher frequency of detections in the surface samples suggests a more recent source of contamination, such as residual from paving of nearby roads and parking lots.

The highest concentrations of PAHs were detected in surface soils collected in the vicinity of paved roadways and/or parking lots in Areas A-1, A-2, and A-3, and may have been a result of asphalt paving activities. High levels were also detected in subsurface samples from a few locations throughout the commercial/industrial properties in Areas A-1 and A-2.

<u>PCBs</u> - Due to their low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, PCBs are more readily adsorbed to soil particles than are either VOC or SVOC contaminants. Furthermore, PCBs present in surface soil have less of a tendency to volatilize into the atmosphere, and PCBs adsorbed to soil particles or present in soil pore spaces have less of a tendency to volatilize into unsaturated soil pore spaces or voids and to potentially migrate to the surface or follow some other preferential flow paths due to their low vapor pressures, compared to the highly volatile VOCs and somewhat volatile SVOCs.

These chemical properties support the frequent detection of elevated levels of PCBs, particularly Aroclor 1262 and Aroclor 1268 (with less frequent detection of Aroclor 1242, Aroclor 1248, Aroclor 1254, and Aroclor 1260), in both surface and subsurface soil throughout the Ferry Creek Area. The high frequency of PCB detection in both surface and subsurface soil suggests that the PCBs were sorbed onto soil particles placed in the area as fill material from the Raymark Facility. Due to the lack of mobility of PCBs in the environment, the PCBs are likely to have remained where they were deposited.

The levels of PCBs were relatively consistent in soil samples throughout the Ferry Creek Area, with the highest levels detected in both surface and subsurface samples north of the culvert under East Broadway in the central and southern regions of Area A-1, south of the culvert/west of the Ferry Creek channel in the northern commercial/industrial properties of Area A-2, and south of the culvert/east of the Ferry Creek channel in the northern region of Area A-3.

<u>Dioxins and Furans (as TEF)</u> - As with PCBs, dioxins and furans have low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, resulting in a tendency for the contaminants to adsorb to soil particles. Furthermore, dioxins and furans present in surface soil have less of a tendency to volatilize into the atmosphere, and those contaminants adsorbed to soil particles or present in soil pore spaces have less of a tendency to volatilize into unsaturated soil pore spaces or voids and to potentially migrate to the surface or follow some other preferential flow paths due to their low vapor pressures, again similar to PCBs.

These chemical properties support the frequent detection of elevated TEF levels in surface soil throughout the Ferry Creek Area, and the somewhat less frequent detection in subsurface soil.

The higher frequency of detections in the surface soils is due to the lack of elevated TEF values in the deeper samples collected. Due to the low likelihood of dioxin/furan mobility in the environment, they are likely to have remained where they were deposited.

The TEF levels were relatively consistent in soil samples throughout the Ferry Creek Area, with the highest levels detected in surface samples north of the culvert under East Broadway in the central region of Area A-1, and south of the culvert and north of the Ferry Creek channel along the eastern border adjacent to wetland area in Area A-3; and subsurface soil north of the culvert in the southern and southwestern regions of Area A-1, south of the culvert and west of the Ferry Creek channel at one of the properties in the northern region of Area A-2, and south of the culvert and north of the Ferry Creek channel along the eastern border adjacent to wetland area in Area A-3.

Metals (primarily Copper and Lead) - Metals tend to be less mobile in the environment than organic contaminants. Under natural conditions, dissolution of metals may occur in the presence of water (due to rainwater runoff or groundwater infiltration) and some metal ions may migrate. However, these dissolved metals can readily precipitate out of solution and adsorb to soil particles. Metals will be more soluble and remain in an aqueous state if the pH is sufficiently low.

These properties support the frequent detection of elevated levels of copper and lead in both surface and subsurface soil throughout the Ferry Creek Area. The somewhat higher frequency of detections in the surface soils is due to the lack of elevated values in the deeper samples collected. Due to the low likelihood of metals mobility in the environment, they are likely to have remained where deposited in the fill material from the Raymark Facility.

The highest levels of both copper and lead were detected in both surface and subsurface samples north of the culvert under East Broadway in region south and west of the Ferry Creek channel in Area A-1, south of the culvert and west of the Ferry Creek channel in the northern properties in Area A-2, and south of the culvert both north and east of the Ferry Creek channel in Area A-3. Elevated concentrations of barium were detected at similar frequencies, with the highest concentrations at the same locations as copper and lead.

It is important to note that elevated levels of lead and/or copper were detected in soil samples submitted for SPLP analysis from regions north of the culvert under East Broadway (including: five soil samples throughout Area A-1) and south of the culvert (including: three soil samples from the commercial/industrial properties in the northern region of Area A-2, and two soil samples in the region north of the Ferry Creek channel in Area A-3). This suggests that these materials may result in a continual release of low levels of copper and lead into the surrounding media.

<u>Asbestos</u> - Asbestos is relatively immobile in the environment, except through bulk advective movement or dispersion in aqueous or airborne environments.

Elevated levels of asbestos were detected relatively frequently in both surface and subsurface soil through the Ferry Creek Area, in similar regions as the presence of elevated levels of copper and lead. Due to the low likelihood of asbestos mobility in the environment, it is likely to have remained where deposited in the fill material from the Raymark Facility.

## E.4.3 Contaminant Fate and Transport in Sediment and Surface Water – Housatonic River Area

Various chemical contaminants are present in the sediment and surface water in the Housatonic River Area. Several pathways are available for contaminants to have entered these media, and for them to subsequently be transported to adjacent media.

The chemical and physical properties of the chemical contaminants affects their ability to migrate into the sediment and surface water, and to subsequently be transported to adjacent media. Contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend adsorb to the sediment particles. These contaminants are apt to remain localized in undisturbed sediment, or to migrate on sediment particulates transported by surface water flows, rainwater runoff, or tidal fluctuations. Contaminants adsorbed to sediment particles may later become desorbed and/or dissolved following contact with solvents or acids, if present. Contaminants with low soil/sediment adsorption coefficients, high water partition coefficients, or high water solubilities would tend to remain in solution and be free to migrate with naturally occurring waters.

Tidal fluctuations may also cause contaminants to be transported into and/or pass through the sediment. Again, those contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend to adsorb to the sediment particles; while those with low soil/sediment adsorption coefficients, high water partition coefficients, and high water solubilities would tend to remain in solution and be free to migrate with natural water sources.

The fate and transport of contamination is, therefore, dependent on the type of chemical contaminant present. A discussion of the fate and transport of the various categories of chemical contaminants in sediment and surface water throughout the Housatonic River Area is presented below. Since no elevated levels of SVOCs (except low levels of phthalates in two samples from Area B) or PCBs were present in surface water samples, and since no samples were collected for analysis of dioxins/furans and asbestos (since these contaminants are virtually insoluble in water), the fate and transport discussion related to surface water is limited to VOCs and metals.

<u>VOCs</u> - Due to their high water solubilities, low soil/sediment partition coefficients, and low octanol-water partition coefficients, VOCs in a sediment/surface water environment tend to remain dissolved in, and free to be transported by, the aqueous media; they do not readily adsorb to sediment particles. Furthermore, VOCs present in surface sediment located in relatively dry locations have more of a tendency to volatilize into the atmosphere than other organic contaminants due to their high vapor pressures. VOCs present in surface water (especially in regions of high turbulence or aeration) also have a tendency to volatilize into the atmosphere due to their high vapor pressures and high air-water partition coefficients.

These chemical properties support the infrequent detection of elevated levels of VOCs (primarily 2-butanone, acetone, and carbon disulfide) in surface sediment, and even less frequent detection in subsurface sediment throughout the Housatonic River Area. VOCs were also detected infrequently in surface water, including: acetone and chlorinated hydrocarbons in a limited number of samples from Area B, and acetone in one sample from Area D; elevated levels of VOCs were not detected in Area C.

The highest concentrations of VOCs were detected in surface sediment from the Ferry Creek channel immediately south of bridge under Ferry Boulevard in the western region, and in the

mouth of the Ferry Creek channel in the northeastern region of the Area B. In both of these areas, factors may have resulted in deposition of sediment, higher-organic content silts, and/or DNAPLs that could encourage adsorption of VOCs onto sediment/silt particles.

SVOCs (primarily PAHs) - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, SVOCs in a sediment/surface water environment are more readily adsorbed to sediment particles than are VOC contaminants. Furthermore, SVOCs present in surface sediment located in relatively dry locations have a tendency to volatilize into the atmosphere to some degree. Their ability to volatilize, however, is less than that of VOCs (with higher vapor pressures) but greater than that of PCBs and dioxins/furans (with much lower vapor pressures).

These chemical properties support the infrequent detection of elevated levels of SVOCs, primarily PAHs, in surface sediment samples and the less frequent detection in subsurface sediment in the Housatonic River Area. The higher frequency of PAH detection in surficial samples suggests that the PAHs were adsorbed onto sediment particles that may have migrated to the areas with surface water flows, rainwater runoff, or tidal fluctuations. Less frequent detections in subsurface sediment suggests that the contaminant source is relatively recent (resulting in surficial deposition of contamination) and/or the particles to which the PAHs adsorb have relatively low densities (resulting in frequent resuspension and surficial redeposition).

The highest concentrations of PAHs were detected in the northwestern region of the Delbuono wetlands adjacent to the lower Ferry Creek channel and downgradient in the channel opposite the wetlands in Area B, along the eastern border adjacent to Shore Road in Area C, and north of the dry dock near the southern portion of the paved parking lot in Area D. These results may have been caused by migration from a nearby source from surface water flows, rainwater runoff, or tidal fluctuations, and subsequent adsorption to sediment particles.

<u>PCBs</u> - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, PCBs in a sediment/surface water environment are more readily adsorbed to sediment particles than are either VOC or SVOC contaminants. Furthermore, PCBs present in surface sediment located in relatively dry locations have little or

no tendency to volatilize into the atmosphere due to their low vapor pressures, compared to the highly volatile VOCs and somewhat volatile SVOCs.

These chemical properties support the relatively frequent detection of elevated levels of PCBs, particularly Aroclor 1262 and Aroclor 1268 (with less frequent detection of Aroclor 016, Aroclor 1242, Aroclor 1248, Aroclor 1254, and Aroclor 1260), in the surface sediment and less frequent detection in subsurface sediment samples collected in the Housatonic River Area. The higher frequency of PCB detection in surficial samples suggests that the PCBs were adsorbed onto sediment particles that may have migrated to the areas with surface water flows, rainwater runoff, or tidal fluctuations. Less frequent detections in subsurface sediment suggests that the contaminant source is relatively recent (resulting in surficial deposition of contamination) and/or the particles to which the PCBs adsorb have relatively low densities (resulting in frequent resuspension and surficial redeposition).

PCBs were detected more frequently in the surface sediment from Ferry Creek channel north of the Delbuono wetlands in Area B, than the two areas further down stream along the Housatonic River. This is believed to be due to the proximity of this area to, and potential contaminant migration from, contaminants present in the Ferry Creek Area. The detection of PCBs in all three areas, however, may also be attributed to the placement of contaminated fill materials from the Raymark Facility in or adjacent to the wetland areas.

<u>Dioxins and Furans (as TEF)</u> - As with PCBs, dioxins and furans have low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, resulting in a tendency for the contaminants to adsorb to sediment particles in a sediment/surface water environment. Furthermore, dioxins and furans present in surface sediment located in relatively dry locations have little or no tendency to volatilize into the atmosphere due to their low vapor pressures, again similar to PCBs.

These chemical properties support the frequent detection of elevated TEF levels in surface sediment and the less frequent detection in subsurface samples throughout the Housatonic River Area. The higher frequency of detection in surface samples suggests that the dioxins and/or furans were adsorbed onto sediment particles that migrated to these areas with surface water flows, rainwater runoff, or tidal fluctuations. Less frequent detections in subsurface

sediment suggests that the contaminant source is relatively recent and/or the particles to which the dioxins and/or furans adsorb have relatively low densities (resulting in frequent resuspension and surficial redeposition).

The highest TEF levels were detected surface sediment in the narrow wetland area in the midwestern region of Area B, both surface and subsurface sediment in the southwestern region and in the wetlands east of Shore Road in Area C, and both surface and subsurface sediment near the dry dock in the southern region of Area D. It is important to note that elevated TEF levels were detected more frequently in the wetland area than in the Ferry Creek channel. These results suggest that the source may be based on surface disposal of contaminated fill materials and/or the migration of dioxins/furans on suspended in surface water flows that tend to accumulate in the wetlands were the sediment/silt media may be higher in organic carbon.

Metals (primarily Copper and Lead) - Metals tend to be less mobile in the environment than organic contaminants. Under natural conditions, dissolution of metals may occur in the presence of water (due to surface water flows, rainwater runoff, or tidal fluctuations) and some metal ions may migrate. However, many of these dissolved metals can readily precipitate out of solution and adsorb to sediment particles. Metals will be more soluble and remain in an aqueous state if the pH is sufficiently low.

These properties support the frequent detection of elevated levels of copper and lead in both surface and subsurface sediment in Area B, the frequent detection in surface sediment and less frequent detection in subsurface sediment in Areas C and D, and a corresponding infrequent detection in surface water (except in the wetlands of Area C).

In the sediment, the similar frequency of detections in surface and subsurface sediment samples in the Ferry Creek channel suggests that deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth) and/or the metals migrated in the dissolved state and precipitated on sediment particles upon contact. Similar detections with depth in the wetland areas adjacent to the lower Ferry Creek channel may be attributable to the placement of contaminated sediments dredged from the mouth of Ferry Creek. In the additional wetland areas further downstream along the Housatonic River, a higher frequency of copper and lead detections in the surface sediment

suggests that the contaminant source is relatively recent (resulting in surficial deposition of contamination) and/or the metals migrated in the dissolved state and precipitated on surface sediment particles upon contact. The distribution of the contaminants suggest the primary source in the placement of contaminated fill materials from the Raymark Facility in or adjacent to these area. Elevated levels of barium were also detected in the same regions, as the copper and lead.

The fact that copper and lead are not apt to dissolve in an aqueous environment under natural conditions (neutral pHs) is supported by their low frequency of detection in surface water throughout Areas B and D. The more frequent detection in surface water from the Area C suggests that these metals may have been present in suspended particulate in this area or were leached out of contaminated sediment or soils. Elevated concentrations of barium (being more water soluble) were detected at slightly higher frequencies in surface water throughout the Housatonic River Area.

It is important to note that elevated levels of lead and/or copper were detected in three sediment samples submitted for SPLP analysis from Area B; no samples were collected for SPLP analysis from the other two areas. This suggests that these materials may result in a continual release of copper and lead into the surrounding media.

<u>Asbestos</u> - Asbestos is relatively immobile in the environment, except through bulk advective movement or dispersion in aqueous or airborne environments.

Elevated levels of asbestos were detected sediment in relatively the same locations as were copper and lead, thereby supporting the conclusions stated previously regarding the placement of contaminated fill material in or adjacent to the wetlands in Areas B, C, and D, as well as the placement of contaminated sediment dredged from the mouth of Ferry Creek in the wetlands adjacent to the lower Ferry Creek channel in Area B.

## E.4.4 Contaminant Fate and Transport in Soil – Housatonic River Area

Various chemical contaminants are also present in the soil located adjacent to the wetlands in the Housatonic River Area. Several pathways are available for contaminants to have entered the soil in these regions, and for them to subsequently be transported to adjacent media.

As with the sediment and surface water, the chemical and physical properties of the chemical contaminants in soil affects their ability to migrate in the Housatonic River Area, and to subsequently be transported to adjacent media. Contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend to adsorb to the soil particles or remain in the soil pore spaces. These contaminants are apt to remain localized in undisturbed soil. Contaminants in soil pore spaces or adsorbed to soil particles may later become mobilized through contact with solvents or acids, precipitation, or groundwater. Contaminants with low soil adsorption coefficients, high water partition coefficients, and high water solubilities would tend to remain in solution and be free to migrate with naturally occurring waters either downward to the water table or laterally to a surface water source.

The fate and transport of contamination is, therefore, dependent on the type of chemical contaminant present. A discussion of the fate and transport of the various categories of chemical contaminants in soil throughout the Housatonic River Area is presented below.

<u>VOCs</u> - Due to their high water solubilities, low soil partition coefficients, and low octanol-water partition coefficients, VOCs are readily leached from soil and transported into an aqueous media. Furthermore, VOCs present in surface soil have more of a tendency to volatilize into the atmosphere, and VOCs absorbed to soil particles or present in soil pore spaces are more apt to volatilize into unsaturated soil pore spaces or voids and may potentially migrate to the surface or follow some other preferential flow paths, than other organic compounds due to their high vapor pressures.

These chemical properties support the infrequent detection of elevated levels of VOCs in surface soil and the somewhat higher frequency in subsurface soil throughout the Housatonic River Area. The higher concentrations of VOCs, primarily carbon disulfide, ketones (2-butanone and acetone), and aromatic hydrocarbons (toluene and xylenes (total)), were

detected in subsurface soils located in a few of locations adjacent to the wetland areas, including: southeast of the Delbuono wetlands in Area B, north and east of the wetlands adjacent to the Housatonic Boat Club in Area C, and in the dry dock area in the southern region of Area D. The presence of these contaminants may have been the result of localized releases at the properties.

SVOCs (primarily PAHs) - Due to their low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, SVOCs are more readily adsorbed to soil particles than are VOC contaminants. Furthermore, SVOCs present in surface soil have a tendency to volatilize into the atmosphere to some degree, and SVOCs adsorbed to soil particles or present in soil pore spaces have a tendency to volatilize into unsaturated soil pore spaces or voids and may potentially migrate to the surface or follow some other preferential flow paths to some degree. Their ability to volatilize, however, is less than that of VOCs (with higher vapor pressures) but greater than that of PCBs and dioxins/furans (with much lower vapor pressures).

These chemical properties support the frequent detection of elevated levels of SVOCs, primarily PAHs, in surface soil and the somewhat lower frequency in subsurface soil throughout the Housatonic River Area. The high frequency of detections suggests that the PAHs were adsorbed onto soil particles placed in the area as fill material. The somewhat higher frequency of detections in the surface samples suggests a more recent source of contamination, such as residual from paving of nearby roads and parking lots.

The highest concentrations of PAHs were detected in surface soils collected in the vicinity of paved roadways and/or parking lots in Areas B, C, and D, and may have been a result of asphalt paving activities.

<u>PCBs</u> - Due to their low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, PCBs are more readily adsorbed to soil particles than are either VOC or SVOC contaminants. Furthermore, PCBs present in surface soil have less of a tendency to volatilize into the atmosphere, and PCBs adsorbed to soil particles or present in soil pore spaces have less of a tendency to volatilize into unsaturated soil pore spaces or voids and to

potentially migrate to the surface or follow some other preferential flow paths due to their low vapor pressures, compared to the highly volatile VOCs and somewhat volatile SVOCs.

These chemical properties support the frequent detection of elevated levels of PCBs, particularly Aroclor 1262 and Aroclor 1268 (with less frequent detection of Aroclor 1254), in both surface and subsurface soil throughout the Ferry Creek Area. The high frequency of PCB detection in both surface and subsurface soil suggests that the PCBs were adsorbed onto soil particles placed in the area as fill material from the Raymark Facility. Due to the lack of mobility of PCBs in the environment, the PCBs are likely to have remained where they were deposited.

The levels of PCBs were relatively consistent in soil samples throughout the Housatonic River Area, with the highest levels detected in subsurface soils west of lowermost extent of the Ferry Creek channel in Area b, and in both surface and subsurface soils both north and east of the wetlands adjacent to the Housatonic Boat Club in Area C, and in the dry dock area in the southern region of Area D.

<u>Dioxins and Furans (as TEF)</u> - As with PCBs, dioxins and furans have low water solubilities, high soil partition coefficients, and high octanol-water partition coefficients, resulting in a tendency for the contaminants to sorb to soil particles. Furthermore, dioxins and furans present in surface soil have less of a tendency to volatilize into the atmosphere, and those contaminants adsorbed to soil particles or present in soil pore spaces have less of a tendency to volatilize into unsaturated soil pore spaces or voids and to potentially migrate to the surface or follow some other preferential flow paths due to their low vapor pressures, again similar to PCBs.

These chemical properties support the frequent detection of elevated TEF levels in the limited number of surface and subsurface soil samples collected throughout the Housatonic River. These contaminants were detected in the same regions as the PCBs and are believed to have been transported to these regions in a similar manner.

Metals (primarily Copper and Lead) - Metals tend to be less mobile in the environment than organic contaminants. Under natural conditions, dissolution of metals may occur in the presence of water (due to rainwater runoff or groundwater infiltration) and some metal ions

may migrate. However, these dissolved metals can readily precipitate out of solution and adsorb to soil particles. Metals will be more soluble and remain in an aqueous state if the pH is sufficiently low.

These properties support the frequent detection of elevated levels of copper and lead in both surface and subsurface soil throughout the Housatonic River Area. The somewhat higher frequency of detections in the surface soils is due to the lack of elevated values in the deeper samples collected. Due to the low likelihood of metals mobility in the environment, they are likely to have remained where deposited in the fill material from the Raymark Facility.

The highest levels of both copper and lead were detected in both surface and subsurface samples located in the southwest of the Delbuono wetlands in Area B, north and east of the wetlands adjacent to the Housatonic Boat Club in Area C, and central and southern regions of Area D. Elevated concentrations of barium were detected at similar frequencies, with the highest concentrations at the same locations as copper and lead.

<u>Asbestos</u> - Asbestos is relatively immobile in the environment, except through bulk advective movement or dispersion in aqueous or airborne environments.

Elevated levels of asbestos were detected infrequently frequently in both surface and subsurface soil throughout the Housatonic River Area, in similar regions as the presence of elevated levels of copper and lead. Due to the low likelihood of asbestos mobility in the environment, it is likely to have remained where deposited in the fill material from the Raymark Facility.

# E.4.5 Contaminant Fate and Transport in Sediment and Surface Water – Elm Street Wetland and Selby Pond Areas

Various chemical contaminants are present in the sediment and surface water in the Elm Street Wetland and Selby Pond Areas. Several pathways are available for contaminants to have entered these media, and for them to subsequently be transported to adjacent media.

The chemical and physical properties of the chemical contaminants affects their ability to migrate into the sediment and surface water, and to subsequently be transported to adjacent media. Contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend adsorb to the sediment particles. These contaminants are apt to remain localized in undisturbed sediment, or to migrate on sediment particulates transported by surface water flows, rainwater runoff, or tidal fluctuations. Contaminants adsorbed to sediment particles may later become desorbed and/or dissolved following contact with solvents or acids, if present. Contaminants with low soil/sediment adsorption coefficients, high water partition coefficients, or high water solubilities would tend to remain in solution and be free to migrate with naturally occurring waters.

Tidal fluctuations may also cause contaminants to be transported into and/or pass through the sediment. Again, those contaminants with high soil/sediment adsorption coefficients and low water partition coefficients would tend to adsorb to the sediment particles; while those with low soil/sediment adsorption coefficients, high water partition coefficients, and high water solubilities would tend to remain in solution and be free to migrate with natural water sources.

The fate and transport of contamination is, therefore, dependent on the type of chemical contaminant present. A discussion of the fate and transport of the various categories of chemical contaminants in sediment and surface water throughout the Elm Street Wetland and Selby Pond Areas is presented below. Since no elevated levels of SVOCs were present in surface water samples, and since no samples were collected for analysis of dioxins/furans and asbestos (since these contaminants are virtually insoluble in water), the fate and transport discussion related to surface water is limited to VOCs, PCBs, and metals.

<u>VOCs</u> - Due to their high water solubilities, low soil/sediment partition coefficients, and low octanol-water partition coefficients, VOCs in a sediment/surface water environment tend to remain dissolved in, and free to be transported by, the aqueous media; they do not readily adsorb to sediment particles. Furthermore, VOCs present in surface sediment located in relatively dry locations have more of a tendency to volatilize into the atmosphere than other organic contaminants due to their high vapor pressures. VOCs present in surface water (especially in regions of high turbulence or aeration) also have a tendency to volatilize into the atmosphere due to their high vapor pressures and high air-water partition coefficients.

These chemical properties support the infrequent detection of elevated levels of VOCs (primarily 2-butanone, acetone, and carbon disulfide) in both surface and subsurface sediment throughout the Elm Street Wetland Area and surface sediment throughout the Selby Pond Area; no subsurface sediment samples were collected for VOC analysis from the Selby Pond Area. In surface water, low levels of acetone and styrene were each detected in a single sample collected in the Elm Street Wetland Area; while, elevated levels of VOCs were not detected in surface water from the Selby Pond Area.

In the sediment, the higher concentrations of VOCs was detected in surface sediment in the wetlands both in the eastern region of the Elm Street Wetland Area and along the eastern side of Selby Pond. This phenomenon may be due to the fact that that these samples were collected close to the outer boundary of the designated wetland areas where surface water was less apt to desorb the contaminants from the sediment.

<u>SVOCs (primarily PAHs)</u> - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, SVOCs in a sediment/surface water environment are more readily adsorbed to sediment particles than are VOC contaminants. Furthermore, SVOCs present in surface sediment located in relatively dry locations have a tendency to volatilize into the atmosphere to some degree. Their ability to volatilize, however, is less than that of VOCs (with higher vapor pressures) but greater than that of PCBs and dioxins/furans (with much lower vapor pressures).

These chemical properties support the detection of elevated levels of SVOCs, primarily PAHs, in sediment samples and the lack thereof in the surface water in the two areas. In the Elm Street Wetland Area, elevated levels of PAHs were detected in only two samples (surface sediment in the eastern region). Infrequent detection of PAHs in the Selby Pond Area were present primarily in surface sediment. The higher frequency of PAH detection in surficial samples, suggests that the PAHs were adsorbed onto sediment particles that may have migrated to these areas with surface water flows, rainwater runoff, or tidal fluctuations. Less frequent detections in subsurface sediment suggest that the contaminant source is relatively recent (resulting in surficial deposition of contamination).

The highest concentrations of PAHs were detected in the northeastern portion of Selby Pond and at two locations in the wetlands along the northeastern and southeastern borders of Selby Pond.

<u>PCBs</u> - Due to their low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, PCBs in a sediment/surface water environment are more readily adsorbed to sediment particles than are either VOC or SVOC contaminants. Furthermore, PCBs present in surface sediment located in relatively dry locations have little or no tendency to volatilize into the atmosphere due to their low vapor pressures, compared to the highly volatile VOCs and somewhat volatile SVOCs.

These chemical properties support the frequent detection of elevated levels of PCBs, particularly Aroclor 1254 (with less frequent detection of Aroclor 1242, Aroclor 1262, and Aroclor 1268), in both the surface and subsurface sediment samples collected in the Elm Street Wetland Area; and the infrequent detection of PCBs, particularly Aroclor 1262 and Aroclor 1268, in the surface sediment and lack of detection in the subsurface sediment in the Selby Pond Area.

The high frequency of PCB detection in the sediment in the Elm Street Wetland Area, localized in the eastern region, suggests that the PCBs were adsorbed onto sediment particles that were placed there as fill material from the Raymark Facility or that may have migrated to this area with surface water flows, rainwater runoff, or through the nearby culvert. The relatively similar frequency of detection in both surface and subsurface sediment deposition of contaminated sediment has occurred over an extended period of time (resulting in similar deposition of contamination with depth) and/or the particles to which the PCBs adsorb have relatively high densities (resulting in infrequent resuspension and redeposition). The highest levels were detected in surface and subsurface sediment in the eastern region of the delineated wetlands and in surface sediment in the northeastern region of the area just beyond the wetland boundary. Moderate levels of Aroclor-1254 were also present in surface water from these areas.

The low frequency of PCB detection in the Selby Pond Area were localized in the surface sediment in the northern region of the pond and in the vicinity of the mouth of the channel

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connecting the pond tidally to the Ferry Creek channel. No PCBs were detected in subsurface sediment or surface water collected in the area.

<u>Dioxins and Furans (as TEF)</u> - As with PCBs, dioxins and furans have low water solubilities, high soil/sediment partition coefficients, and high octanol-water partition coefficients, resulting in a tendency for the contaminants to adsorb to sediment particles in a sediment/surface water environment. Furthermore, dioxins and furans present in surface sediment located in relatively dry locations have little or no tendency to volatilize into the atmosphere due to their low vapor pressures, again similar to PCBs.

These chemical properties support the infrequent detection of elevated TEF levels in surface sediment and the lack of detection in subsurface samples throughout the Elm Street Wetland and Selby Pond Areas.

The TEF levels were fairly consistent in surface samples collected throughout the Selby Pond Area. In the Elm Street Wetland Area, the highest levels were detected in surface sediment from the eastern region of the delineated wetlands. The potential migration pathways are the same as those for PCBs.

Metals (primarily Copper and Lead) - Metals tend to be less mobile in the environment than organic contaminants. Under natural conditions, dissolution of metals may occur in the presence of water (due to surface water flows, rainwater runoff, or tidal fluctuations) and some metal ions may migrate. However, many of these dissolved metals can readily precipitate out of solution and adsorb to sediment particles. Metals will be more soluble and remain in an aqueous state if the pH is sufficiently low.

These properties support the frequent detection of elevated levels of copper and lead in surface sediment and the detection in a single subsurface sediment in the Elm Street Wetland and Selby Pond Areas, the frequent detection in surface water in the eastern region of the Elm Street Wetland, and the detection of a single elevated level of lead in surface water collected in the Selby Pond Area in the vicinity of the mouth of the channel connecting the pond tidally to the Ferry Creek channel.

In the sediment, the higher frequency of detections in surface samples suggests that the contaminant source is relatively recent (resulting in surficial deposition of contamination) and/or the metals migrated in the dissolved state and precipitated on surface sediment particles upon contact. The levels were fairly consistent throughout the Selby Pond Area, with the highest levels detected in the vicinity of the mouth of the channel connecting the pond tidally to the Ferry Creek channel. In the Elm Street Wetland Area, the highest levels were detected in surface sediment from the northern and eastern regions of the delineated wetlands in the vicinity of the culvert; surface water samples from these locations also contained elevated levels of copper and lead.

The fact that copper and lead are not apt to dissolve in an aqueous environment under natural conditions (neutral pHs) is supported by their low frequency of detection in surface water from the Selby Pond Area. The more frequent detection in surface water from the eastern region of the Elm Street Wetlands suggests that these metals may have been present in suspended particulate in this area, leached out of contaminated sediment or soils, or transported to the wetlands through the culvert.

Elevated concentrations of barium were detected at slightly higher frequencies in surface and subsurface sediment, as well as in surface water, in the Elm Street Wetland Area. Barium was also detected at slightly higher frequencies in surface sediment in the Selby Pond Area, but it was not detected in the surface water. The highest concentrations of barium were detected in the same general areas, as copper and lead.

<u>Asbestos</u> - Asbestos is relatively immobile in the environment, except through bulk advective movement or dispersion in aqueous or airborne environments.

Elevated levels of asbestos were detected infrequently in surface sediment from the Selby Pond Area. The highest levels were detected in the northeastern region of the pond in the vicinity of the mouth of the channel connecting the pond tidally to the Ferry Creek channel. The findings suggest that asbestos may have been transported tidally from Ferry Creek, as well as by transport in rainwater runoff from asbestos-laden fill materials placed in close proximity to the north eastern region of Selby Pond.

Asbestos was not detected in surface sediment from the Elm Street Wetland Area. No subsurface samples were collected for asbestos analysis from either area.

#### E.4.6 Contaminant Fate and Transport in Soil – Elm Street Wetland

Low levels of PAHs, copper, and lead were present in the single surface soil sample collected from the western border of the delineated wetlands in the Elm Street Wetland Area. PCBs and dioxins/furans were not detected above background concentrations, and no sample was collected for analysis of VOCs. These results suggest that the soil sample may have been contaminated to a small degree by the placement of fill material from the Raymark Facility, or perhaps more likely to have been the result of rainwater runoff due to the close proximity to Elm Street.

## Appendix F

**Human Health Risk Assessment Supporting Documentation** 

APPENDIX F - Human Health Risk Assessment

Appendix F.1 - Copper and Lead Screening/CLP Comparison

Appendix F.2 - CT Remediation Standard Regulations

Appendix F.3 - Background Concentrations

Appendix F.4 - Comparison to Connecticut Water Quality Standards

Appendix F.5 - Sample Subsets

Appendix F.6 - Statistics and Exposure Point Concentrations

Appendix F.7 - Toxicity Profiles
Appendix F.8 - Sample Calculations

Appendix F.9 - Human Health Risk Assessments Spreadsheets - Area B
Appendix F.10 - Human Health Risk Assessment Spreadsheets - Area C
Appendix F.11 - Human Health Risk Assessments Spreadsheets - Area F

Appendix F.12 - Results of IEUBK Model and TRW Model for Lead

## Appendix F.1

Correlation of Copper and Lead Field Screening Data vs. Fixed Lab Data

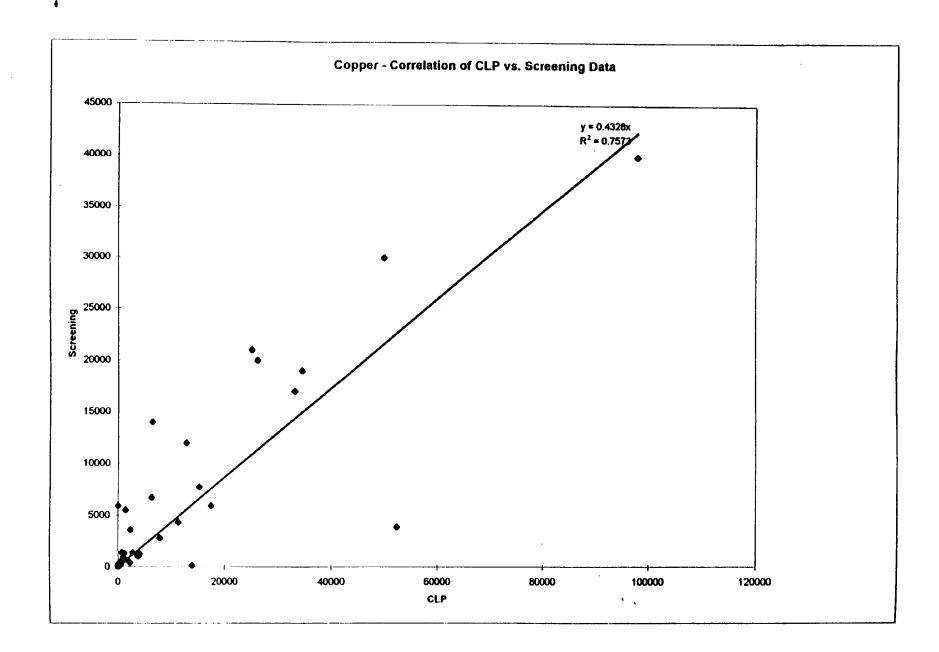
## Appendix F-1 Field Screening – CLP Data Correlation

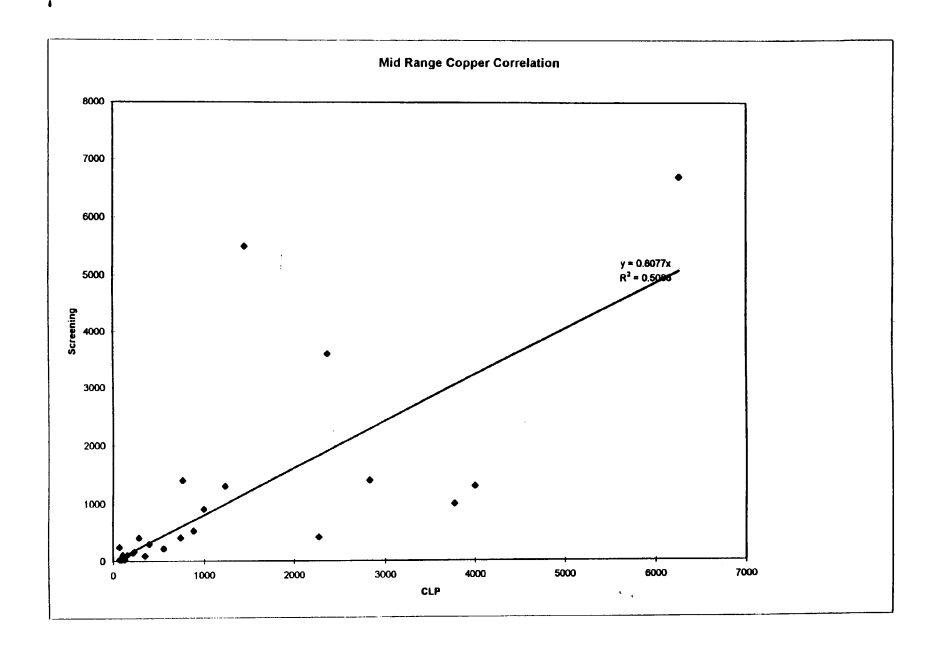
EPA directed Brown and Root Environmental to determine the correlation between data analyzed by field screening and CLP methodologies at the Raymark – Ferry Creek site. A strong correlation would allow for the use of field screening data in quantifying risk at the site. Two statistical procedures were used to determine the correlation between data analyzed by field screening and CLP methodologies: linear regression, which evaluates the correlation on a point-by-point basis; and a nonparametric t-test, which compares the means of two data sets for each method. Paired data selected for the correlation determination were collected at the same location and same depth.

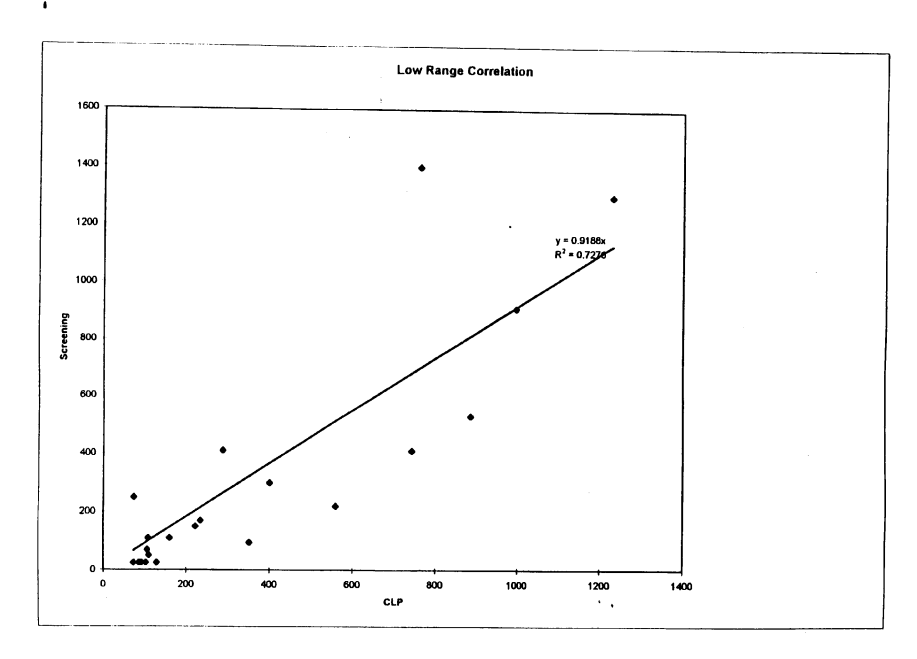
For the first statistical analysis, a scatter plot of paired data was generated for each chemical with the field screening results plotted along the x-axis and the CLP results plotted along the y-axis. A linear regression was then performed on the scatter plot and a correlation coefficient was generated. For data that are strongly correlated, the scatter plot will exhibit a linear relationship with a correlation coefficient (r) of slightly less than 1. The copper and lead data had relatively high correlation coefficients of 0.87 and 0.86, respectively. The PCB data had extremely low correlation coefficients. The PCB scatter plots show that some correlation exists at low concentrations (< 1 ppm), but that this correlation weakens as concentrations increase. This may be due to the narrower calibration ranges of the field screening techniques.

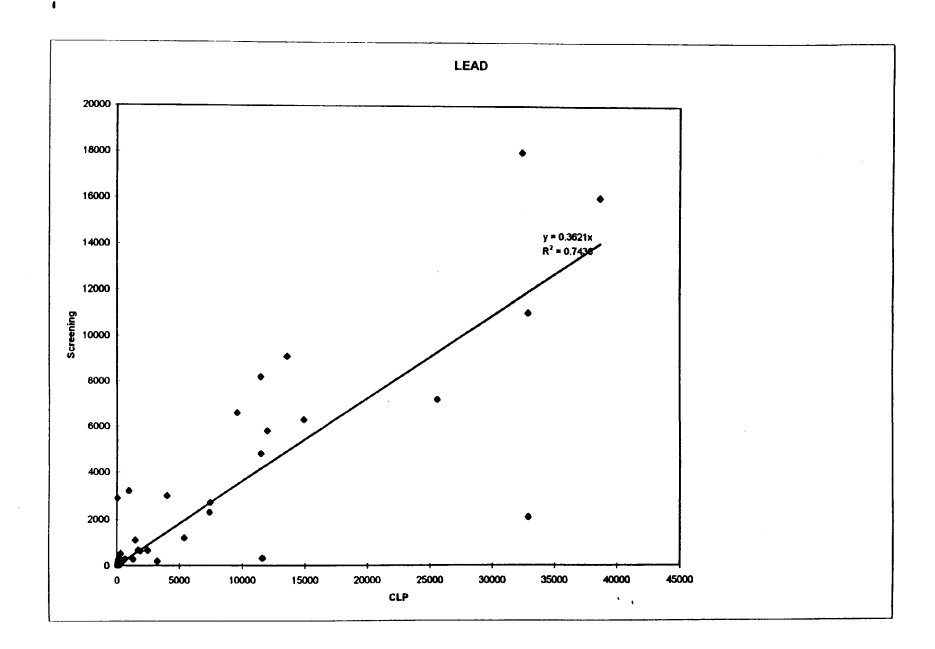
For the second statistical analysis, the field screening data was grouped into one population and the CLP data was grouped into a second population. The Wilcoxon Rank-Sum (WRS) test, a distribution-free or nonparametric t-test, was performed on the two populations to determine whether their means were statistically equivalent. The copper and lead data had statistically equivalent means; the PCB data did not.

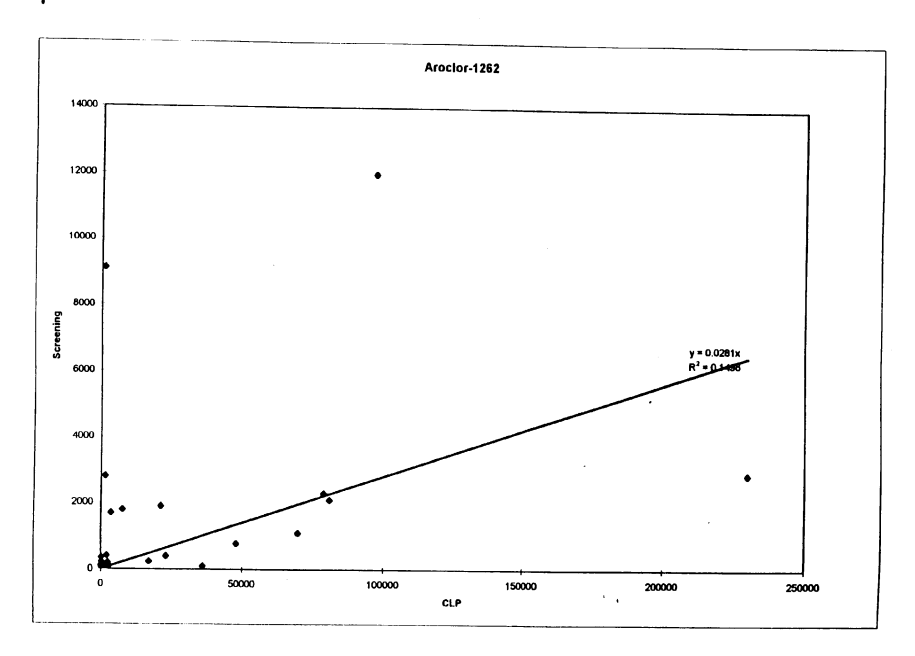
Based on the results of the two statistical analyses, the use of field screening data to quantify risk at the site is acceptable for copper and lead, but not for PCBs



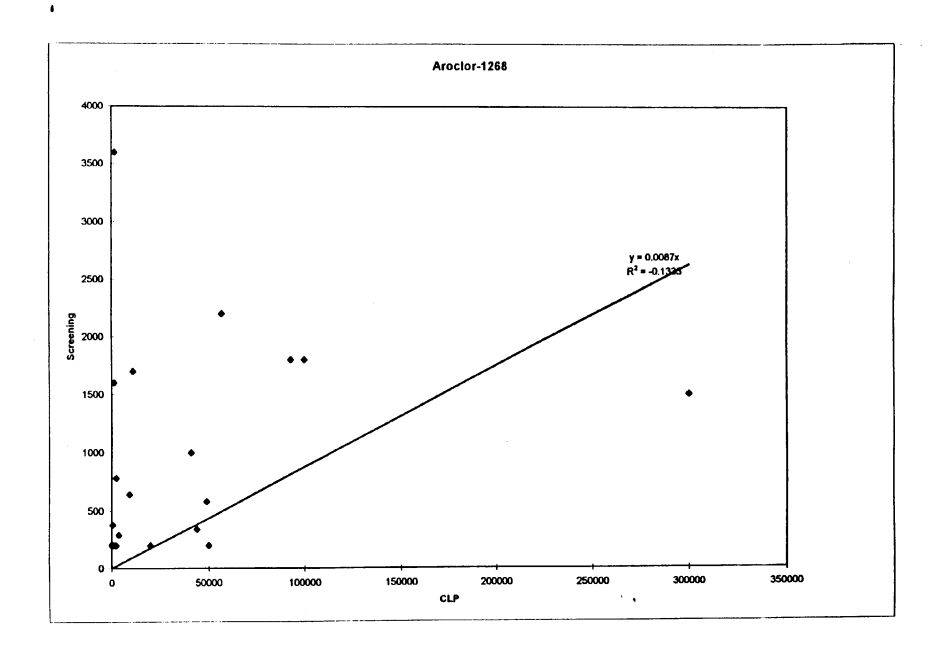












#### Copper Correlation Data

nsample	para	dp_res  dp_res	ult dp	qual ;dp_units	scr_res_scr_res	ult scr qual	dp_res	SCC 100
SP-SO-MW110D-1820	COPPER	7:7		MG/KG	25 ND	U	7	25
MF-SO-MW101D-4850				MG/KG	25 ND	U	7.9	
MF-SO-MW101D-4850				MG/KG	25 ND	<del>- ŭ</del>	8.7	25 25
SP-SO-MW112B-2628	COPPER			MG/KG	25 ND	<del>-</del> Ū	10.6	
MF-SO-MW104D-6062		11.9 11.9	<u> </u>	MG/KG	25 ND	Ŭ		25
MF-SO-MW101D-2830	COPPER		•	MG/KG	25 ND		11.9	25 25
MF-SO-SB5-1416	COPPER	13.8 13.8		MG/KG	8 8	U	12.4	25
MF-SO-MW104D-1618	COPPER			MG/KG	25 ND	<del></del>	13.8	8
MF-SO-MW102-7880	COPPER			MG/KG	25 ND		15.2	25
MF-SO-MW104D-3234	COPPER		<del></del>	MG/KG	25 ND	U	17.2	25 25 25
SP-SO-SB9-0608	COPPER		<del></del>	MG/KG	25 ND	U	17.2	25
MF-SO-MW102-2224	COPPER			MG/KG	25 ND	U	17.4	25
MF-SO-TP3-0405	COPPER	22.1 22.1		MG/KG		U	18.5	25
BC-SO-SB8A-0810	COPPER		'J	MG/KG	53.53	U	22.1	53
MF-SO-MW102-4244	COPPER		_	MG/KG	25 ND	U	22.9	25
MF-SO-MW104D-4648	COPPER				25 ND	Ü	23.4	
SP-SO-SB1-0406B	COPPER		J	MG/KG	25 ND	U .	28.9	25
SP-SO-MW113B-0810	COPPER		<del>- +</del>		21 21	<del></del>	32.9	
MF-SO-MW103-1416	COPPER		J	MG/KG	25 ND	U	33.8	25
MF-SO-SB1-0810	COPPER		<del>-  </del>	MG/KG	5900 5900		34.4	. 5900
BC-SO-SB9-0608	COPPER			MG/KG	26 26	·	35.8	
MF-SO-MW104D-0002	COPPER		J	MG/KG	25 ND	U	37.4	25
SP-SO-SB5-1214	COPPER	63.5 63.5		MG/KG	25 ND	U	40.3	25
SP-SO-SB9-0810	COPPER			MG/KG	25 ND	U	63.5	25
SP-SO-SB7-0204	COPPER		J	MG/KG	250:250	J	73.3	250
BC-SO-SB2-1214	COPPER		J	MG/KG	25 ND	U	73.5	25
SP-SO-MW111D-1012	COPPER		J	MG/KG	25 ND	U	85.4	25
SP-SO-MW110D-0406	COPPER	102!102	J	MG/KG	25 ND	U	91.8	25
MF-SO-MW101D-0608	COPPER	102:102		MG/KG	25 ND	U	102	25
BC-SO-MW120-0406	COPPER		J	MG/KG	70.70	<u>_</u>	106	70
MF-SO-SB2-1416	COPPER	110 110		MG/KG	110 110		108	110
BC-SO-SB8A-1012	COPPER	129:129		MG/KG	51 51		110	51
SP-SO-MW110D-1012	COPPER		J	MG/KG	25 ND	U	129	25
SP-SO-MW111D-0810	COPPER			MG/KG	110:110	J	160	110
BC-SO-SB9-0204B	COPPER		:J	MG/KG		J	222	150
MF-SO-SB8-0608	COPPER	234;234 287;287	J	MG/KG	170:170	J	234	170
MF-SO-SB3-0810	COPPER	267:267		MG/KG	410 410		287	410
SP-SO-SB6-0608A	COPPER	352:352 400:400	J	MG/KG	95,95	·	352.	95
SP-SO-SB3-1416	COPPER			MG/KG	300:300		400	300
MF-SO-SB2-0608	COPPER	5591559	J	MG/KG	220 - 220	·	559:	220
MF-SO-MW103-1618	COPPER			MG/KG	410:410		742	410
SP-SO-MW113B-0406	COPPER	762 762	<del></del>	MG/KG	1400 1400		762	1400
SP-SO-MW113B-0204B	COPPER	885 885	_ <u>J</u>	MG/KG	530 530		885	530
SP-SO-SB8-0002			_ J	MG/KG	910:910		997	910
BC-SO-SB6-0810	COPPER	1230:1230	J	MG/KG	1300:1300		1230	1300
MF-SO-TP2-0506				MG/KG	5500 5500		1450	5500
MF-SO-SB7-1416	COPPER	2270 2270	<u> </u>	MG/KG	420.420		2270	420
SP-SO-MW112B-0810A			<u>J</u>	MG/KG	3600-3600		2370	3600
BC-SO-SB5-0002B	COPPER	283012830		MG/KG	1400 1400		2830	1400
MF-SO-SB7-0406	COPPER	377013770		MG/KG	1000 1000		3770	1000
MF-SO-SB4-1214	COPPER	4000:4000	<u> </u>	MG/KG	1300 1300		4000	1300
MF-SO-SB6-0204	COPPER	6260 6260		MG/KG	6700 6700		6260	6700
SP-SO-MW112B-0608	COPPER	6390 6390		MG/KG	14000 14000		6390	14000
MF-SO-MW102-0406	COPPER	7850 7850		MG/KG	2800 2800	•	7850	2800
SP-SO-S84-0406	COPPER	11300 11300	<u>J</u>	MG/KG	4300 4300		11300	4300
MF-SO-SB4-0406	COPPER	12800 12800	J	MG/KG	12000 12000		12800	12000
	COPPER	13900 13900	_ <u>J</u>	MG/KG	150 150		13900	150
BC-SO-SB3-0204A	COPPER	15200 15200	_ J	MG/KG	7700 7700		15200	7700
BC-SO-SB1-0608	COPPER	17500 17500	J	MG/KG	5900 5900	÷	17500	5900
SP-SO-SB2-0204B	COPPER	25200 25200		MG/KG	21000 21000		25200	21000
BC-SO-SB4-0204	COPPER	26300 26300		MG/KG	20000 20000		26300	20000
BC-SO-SB1-0406	COPPER	33200 33200	J	MG/KG	17000 17000		33200	17000
MF-SO-MW103-0810	COPPER	34600:34600		MG/KG	19000 19000		34600	19000
BC-SO-SB8-0204A	COPPER	49900 49900	J	MG/KG	30000 30000		49900	
							49900	30000

#### Copper Correlation Data

BC-SO-SB1-0204B	COPPER	52300 52300	J	MG/KG	3900 3900	52300	3900
MF-SO-MW103-0608	COPPER	97900 97900		MG/KG	40000 40000	97900	40000

#### Lead Correlation Data

nsample	sample n	o location	рага	clo res	scr_res_clp_qual	scr_qual	units
SP-SO-MW112B-2628	MAES96	SP-SO-MW112B-2628	LEAD	2.1	0:J	U quai	MG/KG
MF-SO-MW104D-3234	MAES95	MF-SO-MW104D-3234	LEAD			<del></del>	MG/KG
MF-SO-MW101D-4850	MAES80	MF-SO-MW101D-4850	LEAD	2.3			
MF-SO-MW101D-4850-0		MF-SO-MW101D-4850	LEAD		0 UJ	U	MG/KG
MF-SO-MW104D-6062	MAET04	MF-SO-MW104D-6062	LEAD			J	MG/KG
SP-SO-MW110D-1820	MAET10	SP-SO-MW110D-1820	LEAD	3		U	MG/KG
MF-SO-MW101D-2830	MAES78	MF-SO-MW101D-2830	LEAD	3.1	0 01		MG/KG
MF-SO-SB5-1416	MAEG35	MF-SO-SB5-1416	LEAD			U	MG/KG
MF-SO-MW102-4244	MAES94	MF-SO-MW102-4244	LEAD	3.8		U	MG/KG
MF-SO-MW104D-4648	MAET03	MF-SO-MW104D-4648	LEAD			U	MG/KG
MF-SO-SB1-0810	MAEG32	MF-SO-SB1-0810	LEAD	5.5	0 J	U	MG/KG
MF-SO-MW102-7880	MAET02	MF-SO-MW102-7880	LEAD	6	10	U	MG/KG
MF-SO-MW102-2224	MAES79	MF-SO-MW102-7680	LEAD	6.5	0.1	U	MG/KG
MF-SO-MW104D-1618	MAES91	MF-SO-MW104D-1618		6.7	0.1	U	MG/KG
BC-SO-SB8A-0810	MAET39	BC-SO-SB8A-0810	LEAD	7.3	0.1	U	MG/KG
MF-SO-MW103-1416	MAES98		LEAD	12.3	0 U	U	MG/KG
BC-SO-SB9-0608	MAET35	MF-SO-MW103-1416 BC-SO-SB9-0608	LEAD	18.2	2900 J		MG/KG
SP-SO-SB1-0406B	MAES73	SP-SO-SB1-0406B	LEAD	20.8	39:	J	MG/KG
MF-SO-MW104D-0002			LEAD	26.6	16 J	<del></del>	MG/KG
SP-SO-SB9-0608	MAES87	MF-SO-MW104D-0002	LEAD	26.8	01	U	MG/KG
SP-SO-MW113B-0810	MAET25	SP-SO-SB9-0608	LEAD	32	0	U	MG/KG
SP-SO-SB5-1214		SP-SO-MW113B-0810	LEAD	40.1	0	U	MG/KG
BC-SO-MW120-0406	MAES83	SP-SO-SB5-1214	LEAD	52.7	0·J	U	MG/KG
MF-SO-TP3-0405	MAES77	BC-SO-MW120-0406	LEAD	55.3	61 J	J	MG/KG
MF-SO-SB2-1416	MAEH50	MF-SO-TP3-0405	LEAD	59.8	173		MG/KG
BC-SO-SB2-1214	MAEG34	MF-SO-SB2-1416	LEAD	72.5	31	·	MG/KG
SP-SO-SB7-0204	MAEF93	BC-SO-SB2-1214	LEAD	75.9	0 J	U	MG/KG
BC-SO-SB8A-1012	MAET06	SP-SO-SB7-0204	LEAD	89.9	53		MG/KG
SP-SO-SB9-0810	MAET40	BC-SO-SB8A-1012	LEAD	94.1	49	<u>J</u>	MG/KG
SP-SO-MW111D-0810-D	MAET38	SP-SO-SB9-0810	LEAD	101	280		MG/KG
MF-SO-MW101D-0608		SP-SO-MW111D-0810	LEAD	122	140		MG/KG
BC-SO-SB9-0204B	MAEH58	MF-SO-MW101D-0608	LEAD	130	71 J		MG/KG
SP-SO-MW1138-0204B	MAET34	BC-SO-SB9-0204B	LEAD		140		MG/KG
SP-SO-MW113B-0406	MAET24	SP-SO-MW113B-0204B		157	120		MG/KG
SP-SO-MW111D-1012	MAET29	SP-SO-MW113B-0406	LEAD	182	170		MG/KG
	MAET23	SP-SO-MW111D-1012	LEAD	183	130		MG/KG
MF-SO-SB3-0810 SP-SO-MW113B-0608	MAEH53	MF-SO-SB3-0810	LEAD	189	18 J		MG/KG
MF-SO-SB3-0810-D	MAET26	SP-SO-MW113B-0608	LEAD	215	210		MG/KG
	MAEH54	MF-SO-SB3-0810	LEAD	227	59 J		MG/KG
SP-SO-SB6-0608A	MAES84	SP-SO-SB6-0608A	LEAD	273	270 J		MG/KG
MF-SO-MW103-1618	MAETOO	MF-SO-MW103-1618	LEAD	275	500 J		MG/KG
SP-SO-MW110D-0406	MAETO8	SP-SO-MW110D-0406	LEAD	305	160 J		MG/KG
MF-SO-SB8-0608	MAET18	MF-SO-SB8-0608	LEAD	310	530		MG/KG
SP-SO-MW111D-0810	MAET21	SP-SO-MW111D-0810	LEAD	354	130		MG/KG
SP-SO-MW110D-1012	MAET13	SP-SO-MW110D-1012	LEAD	363	160 J		MG/KG
SP-SO-SB3-1416	MAES85	SP-SO-SB3-1416	LEAD	459	180 J		MG/KG
MF-SO-SB2-0608	MAEG33	MF-SO-SB2-0608	LEAD	678	290		MG/KG
3C-SO-SB6-0810	MAES76	BC-SO-SEC-3310	LEAD	<del>3</del> 4€	3200 J		MG/KG
MF-SO-TP2-0506	MAEH49	MF-SO-TP2-0506	LEAD	1290	270	. •	MG/KG
SP-SO-SB8-0002	MAET33	SP-SO-SB3-J002	LEAD	1480	1100	-	MG/KG
MF-SO-SB7-0406	MAET17	MF-SO-SB7-0406	LEAD	1690	670	.,	MG/KG
SP-SO-MW112B-0810A	MAES93		LEAD	1870	630 J		MG/KG
BC-SO-SB5-0002B	MAES75	BC-SO-SB5-0002B	LEAD	2460	660 J		MG/KG
MF-SO-SB4-1214	MAEH55	MF-SO-SB4-1214	LEAD	3220	200		MG/KG
MF-SO-SB7-1416	MAET16	MF-SO-SB7-1416	LEAD	3980	3000		MG/KG

#### Lead Correlation Data

SP-SO-MW112B-0608	MAES92	SP-SO-MW112B-0608	LEAD :	53901	1200		140.045
MF-SO-MW102-0406	MAES71				1200 J	<del></del>	MG/KG
		MF-SO-MW102-0406	LEAD	7400	2300 J	1	MG/KG
MF-SO-MW102-0406-D	MAES72	MF-SO-MW102-0406	LEAD	7450	2700 J		MG/KG
MF-SO-SB6-0204	MAEH48	MF-SO-SB6-0204	LEAD .	9600	6600		MG/KG
SP-SO-SB4-0406	MAES86	SP-SO-SB4-0406	LEAD	11500	8200 J	<del></del>	MG/KG
BC-SO-SB1-0608	MAES89	BC-SO-SB1-0608	LEAD	11500	4800 J	<del>i</del>	MG/KG
MF-SO-SB4-0406	MAES88	MF-SO-SB4-0406	LEAD	11600	310 J		MG/KG
BC-SO-SB3-0204A	MAEH57	BC-SO-SB3-0204A	LEAD	12000	5800 J	<del></del>	MG/KG
SP-SO-SB2-0204B	MAES82	SP-SO-SB2-0204B	LEAD	13600	9100iJ	<del></del>	MG/KG
MF-SO-MW103-0810	MAES99	MF-SO-MW103-0810	LEAD	14900	6300;J	<del></del> .i	MG/KG
MF-SO-MW103-0608	MAET01	MF-SO-MW103-0608	LEAD	25600	7200 J		
BC-SO-SB8-0204A	MAET27	BC-SO-SB8-0204A	LEAD	32400	18000		MG/KG
BC-SO-SB4-0204	MAES74	BC-SO-SB4-0204	LEAD	32900	11000 J		MG/KG
BC-SO-SB1-0204B	MAEF91	BC-SO-SB1-0204B	LEAD	32900	2100 J	<del></del>	MG/KG
BC-SO-SB1-0406	MAEF92	BC-SO-SB1-0406	LEAD	38700		<u> </u>	MG/KG
=======================================	1144 (2) 02	100-00-001-0400	LEAD	30700	160001J		MG/KG

#### Aroclor 1262 Correlation Data

nsample	sample_n	o location	para	clp_res sc	r res qual	units	scr_qual
MF-SO-MW104D-6062	SAA687	MF-SO-MW104D-6062	AROCLOR-1262	18.5	100 U	UG/KG	
MF-SO-MW102-2224	SAA663	MF-SO-MW102-2224	AROCLOR-1262		100 U	UG/KG	
MF-SO-MW102-7880	SAA685	MF-SO-MW102-7880	AROCLOR-1262		100 U	UG/KG	
MF-SO-SB5-1416	SA4046	MF-SO-SB5-1416	AROCLOR-1262		100 U	UG/KG	
SP-SO-MW110D-1820	SAA694	SP-SO-MW110D-1820	AROCLOR-1262		100 UJ	UG/KG	
MF-SO-SB1-0810	SA4043	MF-SO-SB1-0810	AROCLOR-1262		100 U	UG/KG	
MF-SO-MW104D-0002	SAA671	MF-SO-MW104D-0002	AROCLOR-1262	20	100 U	UG/KG	
MF-SO-MW104D-3234	SAA678	MF-SO-MW104D-3234	AROCLOR-1262	20	100 U	UG/KG	
MF-SO-MW101D-4850	SAA664	MF-SO-MW101D-4850	AROCLOR-1262		100 U		
MF-SO-MW101D-4850-D	SAA665	MF-SO-MW101D-4850			100 U	UG/KG	
MF-SO-MW101D-2830	SAA662	MF-SO-MW101D-2830	AROCLOR-1262			UG/KG	
SP-SO-MW112B-2628	SAA679	SP-SO-MW112B-2628	AROCLOR-1262		100 U	UG/KG	<u>. U</u>
MF-SO-MW 102-4244	SAA677	MF-SO-MW102-4244	AROCLOR-1262		100 U	UG/KG	
MF-SO-MW104D-4648	SAA686	MF-SO-MW104D-4648		22.5	100 U	UG/KG	
SP-SO-SB1-0406B	SAA651	SP-SO-SB1-0406B	AROCLOR-1262		100 U	UG/KG	
BC-SO-SB9-0608	SA9041	BC-SO-SB9-0608	AROCLOR-1262	26	100 J	UG/KG	
MF-SO-MW104D-1618	SAA674		AROCLOR-1262	29	100 J	UG/KG	
MF-SO-MW103-1416	SAA681	MF-SO-MW104D-1618		31.5	100 U	UG/KG	
SP-SO-SB5-1214	SAA667	MF-SO-MW103-1416	AROCLOR-1262	38.5	100 U	UG/KG	
SP-SO-MW113B-0810		SP-SO-SB5-1214	AROCLOR-1262	51	100 J	UG/KG	
BC-SO-SB8A-0810	SA2742	SP-SO-MW113B-0810	AROCLOR-1262	58	100 J	UG/KG	U
SP-SO-MW113B-0204B	SA2773	BC-SO-SB8A-0810	AROCLOR-1262	69	100 J	UG/KG	
MF-SO-SB2-1416	SA2741	SP-SO-MW113B-0204B		72	100 J	UG/KG	U
	SA4045	MF-SO-SB2-1416	AROCLOR-1262	76	100 J	UG/KG	U
MF-SO-TP3-0405	SAA640	MF-SO-TP3-0405	AROCLOR-1262	79	100	UG/KG	U
SP-SO-SB7-0204	SA2748	SP-SO-SB7-0204	AROCLOR-1262	90	100 J	UG/KG	U
SP-SO-MW113B-0406	SA2746	SP-SO-MW113B-0406	AROCLOR-1262	99	10 <b>0</b> J	UG/KG	U
SP-SO-MW110D-0002	SAA696	SP-SO-MW110D-0002	AROCLOR-1262		100 J	UG/KG	U
BC-SO-SB9-0204B	SAA656	BC-SO-SB9-0204B	AROCLOR-1262	180	100 J	UG/KG	U
SP-SO-SB9-0810	SA9043	SP-SO-SB9-0810	AROCLOR-1262	230	360 J	UG/KG	
MF-SO-SB3-0810	SAA652	MF-SO-SB3-0810	AROCLOR-1262	230	100 J	UG/KG	U
SP-SO-MW110D-0406	SAA692	SP-SO-MW110D-0406	AROCLOR-1262	240	100 J	UG/KG	ับ
SP-SO-SB6-0608A	SAA668	SP-SO-SB6-0608A	AROCLOR-1262	260	100 J	UG/KG	U
MF-SO-SB4-1214	SAA654	MF-SO-SB4-1214	AROCLOR-1262	280	100	UG/KG	
BC-SO-SB2-1214	SAA645	BC-SO-SB2-1214	AROCLOR-1262	280	220	UG/KG	
MF-SO-SB3-0810-D	SAA653	MF-SO-SB3-0810	AROCLOR-1262	320	100 J	UG/KG	U
BC-SO-SB8A-1012	SA2774	BC-SO-SB8A-1012	AROCLOR-1262	330	100 J	UG/KG	Ū
SP-SO-MW111D-0810-D		SP-SO-MW111D-0810	AROCLOR-1262	340	100 J	UG/KG	U
MF-SO-SB8-0608	SA2734	MF-SO-SB8-0608	AROCLOR-1262	340	100 J	UG/KG	
	SAA661	BC-SO-MW120-0406	AROCLOR-1262	380	100	UG/KG	
SP-SO-MW111D-0810	SA2738	SP-SO-MW111D-0810	AROCLOR-1262	500	100 J	UG/KG	
MF-SO-MW101D-0608	SAA648	MF-SO-MW101D-0608	AROCLOR-1262	540	100	UG/KG	
MF-SO-TP2-0506	SA4048	MF-SO-TP2-0506	AROCLOR-1262	550	100	UG/KG	
SP-SO-SB9-0608	SA9042	SP-SO-SB9-0608	AROCLOR-1262	580	100 J	UG/KG	
MF-SO-SB7-0406	SA2733	MF-SO-SB7-0406	AROCLOR-1262	650	100 J	UG/KG	
SP-SO-SB4-0406	SAA670	SP-SO-SB4-0406	AROCLOR-1262		9100	UG/KG	···
MF-SO-SB2-0608	SA4044	MF-SO-SB2-0608	AROCLOR-1262		2800 J	UG/KG	
BC-SO-SB5-0002B	SAA659	BC-SO-SB5-0002B	AROCLOR-1262	2100	410 J	UG/KG	
MF-SO-MW102-0406-D	SAA650	MF-SO-MW102-0406	AROCLOR-1262	2500	200	UG/KG	
SP-SO-SB3-1416	SAA669	SP-SO-SB3-1416	AROCLOR-1262	2800	100 J	UG/KG	
MF-SO-MW 102-0406	SAA649	MF-SO-MW102-0406	AROCLOR-1262	2900	100 3	UG/KG	
:	SAA675	SP-SO-MW112B-0608	AROCLOR-1262		1700 J		U
<u></u>	SAA655	SP-SO-SB8-0002	AROCLOR-1262			UG/KG	Ì
	SAA643	BC-SO-SB1-0204B	AROCLOR-1262	17000	1800 J	UG/KG	ļ
:	SAA682	MF-SO-MW103-0810	AROCLOR-1262			UG/KG	
=	SAA660	BC-SO-SB6-0810	AROCLOR-1262		1900	UG/KG	. [
	SAA690	MF-SO-SB7-1416		23000	400	UG/KG	
	0. 0.000	14117	AROCLOR-1262	36000	100 J *	UG/KG	U

#### Aroclor 1262 Correlation Data

MF-SO-S86-0204	SA4047	MF-SO-SB6-0204	AROCLOR-1262	48000	780	UG/KG
BC-SO-SB3-0204A	SAA647	BC-SO-SB3-0204A	AROCLOR-1262	70000	1100	UG/KG
BC-SO-SB4-0204	SAA658	BC-SO-SB4-0204	AROCLOR-1262	79000	2300	UG/KG
BC-SO-SB1-0406	SAA644	BC-SO-SB1-0406	AROCLOR-1262		2100	UG/KG
SP-SO-SB2-0204B	SAA666	SP-SO-SB2-0204B	AROCLOR-1262		12000	UG/KG
BC-SO-SB8-0204A	SA2744	BC-SO-SB8-0204A	AROCLOR-1262		2900 J	UG/KG

#### Aroclor 1268 Correlation Data

nsample		no location	para	clp res	scr_res cin	qual scr_qua	l units
MF-SO-MW102-2224	SAA663	MF-SO-MW102-2224	AROCLOR-1268	3		U Qua	UG/KG
MF-SO-MW102-7880	SAA685	MF-SO-MW102-7880	AROCLOR-1268	3.5		Ü	UG/KG
MF-SO-TP3-0405	SAA640	MF-SO-TP3-0405	AROCLOR-1268			- U	UG/KG
SP-SO-SB1-0406B	SAA651	SP-SO-SB1-0406B	AROCLOR-1268			- Ū	UG/KG
BC-SO-SB9-0608	SA9041	BC-SO-SB9-0608	AROCLOR-1268	33		<del></del>	UG/KG
MF-SO-MW103-1416	SAA681	MF-SO-MW103-1416	AROCLOR-1268			<del></del> -	UG/KG
MF-SO-MW104D-6062	SAA687	MF-SO-MW104D-6062	AROCLOR-1268			<del>U</del>	
MF-SO-SB5-1416	SA4046	MF-SO-SB5-1416	AROCLOR-1268			<del></del>	UG/KG
SP-SO-MW110D-1820	SAA694	SP-SO-MW110D-1820	AROCLOR-1268			<del>- U</del>	UG/KG
MF-SO-SB1-0810	SA4043	MF-SO-SB1-0810	AROCLOR-1268				UG/KG
MF-SO-MW104D-0002	SAA671	MF-SO-MW104D-0002	AROCLOR-1268			<u>U</u>	UG/KG
MF-SO-MW104D-3234	SAA678	MF-SO-MW104D-3234	AROCLOR-1268			U	UG/KG
MF-SO-MW101D-4850	SAA664	MF-SO-MW101D-4850	AROCLOR-1268			U	UG/KG
MF-SO-MW101D-4850-0		MF-SO-MW101D-4850	AROCLOR-1268			U	UG/KG
MF-SO-MW101D-2830	SAA662	MF-SO-MW101D-2830	AROCLOR-1268			U	UG/KG
SP-SO-MW112B-2628	SAA679	SP-SO-MW112B-2628	AROCLOR-1268		200 U	U	UG/KG
MF-SO-MW102-4244	SAA677	MF-SO-MW102-4244			200·U	U	UG/KG
MF-SO-MW104D-4648	SAA686	MF-SO-MW104D-4648	AROCLOR-1268			U	UG/KG
SP-SO-SB6-0608A	SAA668	SP-SO-SB6-0608A	AROCLOR-1268		200 U	U	⊎G/KG
MF-SO-MW104D-1618	SAA674		AROCLOR-1268		200 J	U	"UG/KG
SP-SO-MW113B-0204B		MF-SO-MW104D-1618	AROCLOR-1268	63	200 U	U	UG/KG
SP-SO-MW113B-0810	SA2741	SP-SO-MW113B-0204B			200 J	U	UG/KG
BC-SO-SB8A-0810	SA2742	SP-SO-MW113B-0810	AROCLOR-1268		200 J	U	'UG/KG
SP-SO-SB5-1214		BC-SO-SB8A-0810	AROCLOR-1268		200 J	U	UG/KG
MF-SO-SB2-1416	SAA667	SP-SO-SB5-1214	AROCLOR-1268	84	200	U	UG/KG
BC-SO-MW120-0406	SA4045	MF-SO-SB2-1416	AROCLOR-1268	92	200	U	UG/KG
SP-SO-SB7-0204	SAA661	BC-SO-MW120-0406	AROCLOR-1268	120	200 J	U	UG/KG
BC-SO-SB9-0204B	SA2748	SP-SO-SB7-0204	AROCLOR-1268	130	200 J	U	UG/KG
	SAA656	BC-SO-SB9-0204B	AROCLOR-1268	130	200 J	U	UG/KG
SP-SO-MW113B-0406	SA2746	SP-SO-MW113B-0406	AROCLOR-1268	150	200 J	U	UG/KG
SP-SO-MW110D-0002 MF-SO-SB3-0810	SAA696	SP-SO-MW110D-0002	AROCLOR-1268	200	210 J		UG/KG
SP-SO-SB3-0810 SP-SO-SB9-0608	SAA652	MF-SO-SB3-0810	AROCLOR-1268	250	200 J	U	UG/KG
MF-SO-SB4-1214	SA9042	SP-SO-SB9-0608	AROCLOR-1268	260	200 J	U	UG/KG
	SAA654	MF-SO-SB4-1214	AROCLOR-1268	260	200	U	UG/KG
SP-SO-MW110D-0406	SAA692	SP-SO-MW110D-0406	AROCLOR-1268	270	200 J	U	UG/KG
MF-SO-SB7-0406	SA2733	MF-SO-SB7-0406	AROCLOR-1268	340	200 J	U	UG/KG
BC-SO-SB2-1214	SAA645	BC-SO-SB2-1214	AROCLOR-1268	340	200	. U	UG/KG
MF-SO-SB3-0810-D SP-SO-SB9-0810	SAA653	MF-SO-SB3-0810	AROCLOR-1268	350	200 J	Ū ==	UG/KG
	SA9043	SP-SO-SB9-0810	AROCLOR-1268	400	380 J		UG/KG
SP-SO-MW111D-0810	SA2738	SP-SO-MW111D-0810	AROCLOR-1268	420	200 J	U	UG/KG
	SA2774	BC-SO-SB8A-1012	AROCLOR-1268	490	200 J		.UG/KG
<del></del>	SA2734	MF-SO-SB8-0608	AROCLOR-1268	490	200 J	U	UG/KG
MF-SO-MW101D-0608	SA4048	MF-SO-TP2-0506	AROCLOR-1268	500	200	U	UG/KG
NF-SU-MVV 101D-0608	SAA648	MF-SO-MW101D-0608	AROCLOR-1268	620	200	U	UG/KG
SP-SO-MW111D-0810-D		SP-SO-MW111D-0810	AROCLOR-1268	760	200 J	U	UG/KG
MF-SO-SB2-0608	SA4044	MF-SO-SB2-0608	AROCLOR-1268	1200	1600		UG/KG
SP-SO-SB4-0406	SAA670	SP-SO-SB4-0406	AROCLOR-1268	1600	3600		UG/KG
	SAA650	MF-SO-MW102-0406	AROCLOR-1268	1600	200	U	UG/KG
	SAA649	MF-SO-MW102-0406	AROCLOR-1268	1800	200	U	UG/KG
SP-SO-SB3-1416	SAA669	SP-SO-SB3-1416	AROCLOR-1268	2400	200	U	UG/KG
	SAA675	SP-SO-MW112B-0608	AROCLOR-1268	2400	780		UG/KG
TIT 2 15	SAA659	BC-SO-SB5-0002B	AROCLOR-1268	3800	290	- •	UG/KG
	SAA682	MF-SO-MW103-0810	AROCLOR-1268	9300	640		UG/KG
	SAA655	SP-SO-SB8-0002	AROCLOR-1268	11000	1700 J		UG/KG
*_ ** _ * . * . * . *	SAA643	BC-SO-SB1-0204B	AROCLOR-1268	20000	200	··- ·· ·· · · · · · · · · · · · · · · ·	UG/KG
	SAA660	BC-SO-S86-0810	AROCLOR-1268	41000	1000		UG/KG
C-SO-SB3-0204A	SAA647	BC-SO-SB3-0204A	AROCLOR-1268	44000	340	• • • • • • • • • • • • • • • • • • • •	UG/KG
	SA4047	MF-SO-SB6-0204	AROCLOR-1268	49000	580		UG/KG
IF-SO-SB7-1416	SAA690	MF-SO-SB7-1416	AROCLOR-1268	50000	200 J *	U	UG/KG
							33,10

#### Aroclor 1268 Correlation Data

SP-SO-SB2-0204B	SAA666	SP-SO-SB2-0204B	AROCLOR-1268: 57000	2200	I IIICIKO
BC-SO-SB4-0204	SAA658	BC-SO-SB4-0204	AROCLOR-1268 93000		UG/KG
BC-SO-SB1-0406	SAA644	BC-SO-SB1-0406		1800 i	UG/KG
BC-SO-SB8-0204A			AROCLOR-1268 100000	1800:	UG/KG
DC-30-388-0204A	SA2744	BC-SO-SB8-0204A	AROCLOR-1268 300000	1500 J	UG/KG

## Lead Linear Regression Output

#### **SUMMARY OUTPUT**

Regression Statistics						
Multiple R	0.864407719					
R Square	0.747200705					
Adjusted R Square	0.743427582					
Standard Error	4504.308087					
Observations	69					

	df		SS	MS	F	Significance F
Regression		1	4017837739	4017837739	198.032385	1.10015E-21
Residual		67	1359349020	20288791.34		
Total		68	5377186759			

	Coefficients	Standard Error	t Stat	P-value	Lower 95%	Upper 95%	Lower 50.0%	Upper 50 0%
Intercept	557.588161	600.8487517	0.928000865	0.356738283	-641.7112978		150.1110006	965.0653215
X Variable 1	2.124784321				1.823407937			

## Copper Linear Regi on Output (Low-range Data)

#### SUMMARY OUTPUT

Regression Statistics							
Multiple R	0.852977736						
R Square	0.727571018						
Adjusted R Square	0.677571018						
Standard Error	212.3416465						
Observations	21						

	d <b>f</b>	SS	MS	F	Significance F
Regression Residual	2	1 2408365.741	2408365.741 45088.97484		6.24751E-07
Total	2	1 3310145.238	45088.97484		

1-4	Coefficients	Standard Error	t Stat	P-value	Lower 95%	Upper 95%	Lower 95.0%	Upper 95.0%
Intercept X Variable 1	0.918 <b>791541</b>	#N/A 0.092076695	#N/A	#N/A	#N/A	#N/A	#N/A	#N/A
	0.010101011	0.032070033	9.978546048	3.28009E-09	0.72672301	1.110860072	0.72672301	1.110860072

## Copper Linear Regression Output (Mid-range Data)

#### SUMMARY OUTPUT

Regression Statistics					
Multiple R	0.713327397				
R Square	0.508835975				
Adjusted R Square	0.471798938				
Standard Error	1153.781792				
Observations	28				

Pogranian	df	SS	MS	F	Significance F
Regression Residual Total	1 27 28	37235945.5 35942735.46 73178680.96		27.9714528	1.57064E-05

Interest	Coefficients	Standard Error	t Stat	P-value	Lower 95%	Upper 95%	Lower OF 00/	1105-00/
Intercept X Variable 1	0.807749021	#N/A 0.118003473	#N/A 6.845120220	#N/A	#N/A	#N/A	Lower 95.0% #N/A	<i>Upper 95.0%</i> #N/A
		31110000110	0.040123233	2.3605E-07	0.565626057	1.049871986	0.565626057	1.049871986

#### SUMMARY OUTPUT

Regression Statistics				
Multiple R	0.870217975			
R Square	0.757279324			
Adjusted R Square	0.741654324			
Standard Error	3786.611241			
Observations	65			

			MS		Significance F
Regression Residual Total	1 64 65	2863061909 917659180.1 3780721089	2863061909 14338424.69	199.6775776	3.35283E-21

Int.	Coefficients	Standard Error	t Stat	P-value	Lower 95%	Upper 95%	Lower 95.0%	Upper 95.0%
Intercept X Variable 1	0	#N/A	#N/A	#N/A	#N/A	#N/A	#N/A	#N/A
A Variable 1	0.432816293	0.027008707	16.02506504	4.38521E-24	0.378860246			0.48677234

## Aroclor 1262 Linear Regression Output

## SUMMARY OUTPUT

Regression S	Statistics
Multiple R	0.386977446
R Square	0.149751544
Adjusted R Square	0.133358101
Standard Error	1802.075377
Observations	62

Total	SS 34890018.25 198096015.6 232986033.9	MS 34890018.25 3247475.666	10.74373509	Significance F 0.001742955
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4.001029303 2.3955E-05 0.015775746 0.040498888 0.015775746 0.040498888
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## SUMMARY OUTPUT

Regression S	Statistics
Multiple R	0.429221558
R Square	0.184231146
Adjusted R Square	0.170634998
Standard Error	38817.30776
Observations	62

Residual 60 90407002918 1506783382 61 1.10824E+11	Pograneia	df	SS	MS	F	Significance F
		60	90407002918		13.55024609	

X Variable 1 29.09476683 7.90389732 3.681065891 0.00499429 13.38463337 4.0014277 -4326.106874 3916.14193	Intercent	Coefficients	Standard Error	t Stat	P-value	Lower 95%	Upper 95%	1 50 001	
10.20 102001 TT,0043 1023 7.3 1.3 1 1/hd / 3d Abb 45 / 1	Intercept X Variable 1	-204.9824707 29.09476683			0.973185972	-12352.73494		-4326.106874	Upper 50.0% 3916.141933 34.45835719

## Copper t-Test Output

# t-Test: Two-Sample Assuming Equal Variances

Mean	Variable 1	Variable 2
Variance	6700.495385	3549.830769
Observations	261525929.3	59073767.02
Pooled Variance	65	65
Hypothesized Mean Difference	160299848.2	
df	0	
t Stat	128	
P(T<=t) one-tail	1.418656658	
Critical one-tail	0.079214719	
P(T<=t) two-tail	1.656844688	
Critical two-tail	0.158429439	
di (Wo-tali	1.978669388	

## Lead t-Test Output

t-Test: Two-Sample Assuming Equal Variances

Mean	Variable 1	Variable 2
	4199.56087	1714.043478
Variance	79076275.86	13087412.01
Observations	69	69
Pooled Variance	46081843.94	09
Hypothesized Mean Difference df	0	
t Stat	136	
	2.150608854	
P(T<=t) one-tail	0.016637564	
t Critical one-tail	1.656135282	
P(T<=t) two-tail	0.033275128	
t Critical two-tail	1.977559805	

Appendix F.2

Summary of RSRs Developed by B&RE for New London (1998)

.4121921.7791 FAX: (412) 921-4-4;

C-49-12-7-188

December 23, 1997

Brown & Root Environmental Project Number 7237

Mr. Mark Lewis
Connecticut Department of Environmental Protection
Water Management Bureau
Permitting, Enforcement, and Remediation Division
Federal Remediation Program
79 Elm Street
Hartford, Connecticut 06106-5127

Reference:

CLEAN Contract No. N62472-90-D-1298

Contract Task Order No. 0260

Subject:

Calculated CTDEP Remediation Standards
Lower Subase Remedial Investigation

Naval Submanne Base - New London, Groton, Connecticut

#### Dear Mr. Lewis:

in preparation of the Lower Subase Remedial Investigation (RI) Report. Brown & Root (B&R) Environmental has calculated Remediation Standards following the State of Connecticut Remediation Standard Regulations of January 1996. Standards were developed for all chemicals that were analyzed for during the RI sampling and analysis program that did not have previously established CTDEP standards. The intent of this memo is to identify the sources of the standards to be used in the RI Report and to identify those values which have been developed by B&R Environmental using the State guidance.

Background information and the calculated Remediation Standards are provided in Table 1 and Table 2, respectively, which are enclosed. Table 1 summanzes the basis for the chemical-specific remediation standards (i.e., promulgated, calculated, or calculated using a sumogate) to be included in the RI. The calculated soil Direct Exposure and Pollutant Mobility standards as well as the Groundwater Standard, are provided in Table 2.

It should be noted that pollutant mobility and groundwater standards for GA classified groundwater are provided in Table 2 for completeness. The groundwater at the Lower Subase is classified as GB, therefore standards applicable to GB classified groundwater will be emphasized in the Lower Subase RI Report.



Mr. Mark Lewis

Connecticut Department of Environmental Protection
December 23, 1997 - Page 2

8&R Environmental intends on using these criteria, as well as Region III RBCs, as part of the human health risk assessment for the Lower Subase RI to screen for chemicals of potential concern. Therefore, 8&R Environmental, on the behalf of the United States Navy, requests that the CTDEP review and approve the standards in Table 2. It is hoped that prior approval of the criteria will alleviate unnecessary revisions to the RI Report in the future and expedite any additional risk-related work required by the State (i.e., application for use of alternative criteria).

Due to the current time constraints for preparing the Lower Subase RI, it is requested that the CTDEP complete their review by no later than January 16, 1997. If you have any questions regarding the information provided in the tables or the schedule for the review please contact Mr. Mark Evans at (610) 595-0567 (ext. 162) or me at (412) 921-8244.

Very truly yours.

Corgo A. Rich; P.E. Project Manager

Enclosure(s)

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Mr. Mark Evans, NORTHDIV

Mr. Richard Conant, NSB-NLON Environmental

Ms. Karen Smecker, B&R Environmental

File: CTO 0260

#### TABLE 1

# SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 1 OF 4

			Basis of Value to be Used in RI Report			
Chemical	CAS	Chemical	Promuigated	Calculated	Surrogate	
	Number	Fraction	Value <sup>(1)</sup>	Value <sup>(2)</sup>	Calculated Value <sup>(3)</sup>	
Acenaphthene	83329	SVOC		X		
Acenaonthylene	208968	SVOC	Х			
Anthracene	120127	SVOC	X			
Acetone	67641	VOC	Х			
Aldrin	309002	PEST		×		
Aluminum	7429905	INORG	(4)	(4)	(4)	
Antimony	7440360	INORG	Х		<del></del>	
Arsenic	7440382	INORG	X		<del>'</del>	
Banum	7440393		X			
Benzene	71432		X		<del></del>	
Benzia)anthracene	56 <b>553</b>		X		<del></del>	
Benzo(b)fluoranthene	205992		X		<del></del>	
Benzo(k)fluoranthene	207089	SVOC	X		<del>                                     </del>	
Benzo(g,h,i)perylene	191242	SVOC			X	
					(naphthalene)	
Benzo(a)pyrene	50328	SVOC	X	· · · · · · · · · · · · · · · · · · ·		
Beryllium	7440417	INORG	X			
BCH (alpha-)	319846	PEST		X		
BCH (beta-)	319857	PEST		Х		
BCH (delta-)	319868	PEST		· · · · · · · · · · · · · · · · · · ·	X	
	1				(alpha-8HC)	
BCH (gamma-: Lindane)	58899	PEST	X			
Bis(2-chloroethoxy)methane	111911	svoc	(5)	,5)	(5)	
Bis(2-chloroethyl)ether	111444	SVOC	X			
Bis(2-ethylhexyl)phthalate	117817	SVOC	X			
Bromochloromethane	74975	voc			X	
] L		1	1		(chloromethane)	
Bromodichioromethane	75274	VOC	!	X		
Bromoform	75 <b>252</b>	voc	X			
Bromomethane	74839	voc		X		
4-Bromophenyl-phenylether	101553	SVOC		Х		
2-Butanone	78933	VOC	X			
Butylbenzylphthalate	85687	SVOC	X			
Cadmium	7440439	INORG	X			
Calcium	7440702	INORG	(ä)	Ġ)	(ô)	
Carbazole	86748	SVOC		X	1	
Carbon disulfide	75150	VOC	i	X	ī	
Carbon tetrachionide	56235	VOC	X		1	
Chiordane (alpha-)	57749	PEST	X <sup>(i)</sup>		<del></del>	
Chlorgane (gamma-)	57749	PEST	X <sup>(7)</sup>		<u> </u>	
4-Chloroaniline	106478	SVOC		×	<del> </del>	
Chloropenzene	108907	VOC	<u> </u>			
Chlorodibromomethane	124481	VOC	- <del>x</del>		1	
Chloroethane	75003	VOC		X	<u> </u>	
Chloroform	67663	VOC	x		<u> </u>	
Chloromethane	74873	VOC		X	<u> </u>	

#### TABLE 1

# SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 2 OF 4

<b>_</b>		<u>.</u>	Basis of Value to be Used in RI Report			
Chemical	CAS Number	Chemical Fraction	Primulgated alue <sup>(1)</sup>	Calculated  Value <sup>(2)</sup>	Surrogate Calculated Value <sup>(3)</sup>	
4-Chioro-3-methylphenoi	59507	SVOC			X (3-methylphenol	
2-Chloronaphthalene	91 <b>587</b>	SVOC I		X	1	
2-Chiorophenol	95578		X			
4-Chloropnenyi-phenylether	7005723	SVOC			X (4-Bromopheny) phenylether)	
Chromium (total)		INORG	X(6)			
Chrysene	218019	SVOC		X		
Cobart	7440484	INORG		X		
Copper	7440508	INORG	(4)	(4)	(4)	
4.4'-ODD	72548			X		
4,4'-ODE	72559			Х		
4.4'-DDT	50293			X		
Dibenzofuran	132649			Х		
Oibenz(a,h)anthracene	53703	SVOC		X		
1,2-Dibromo-3-chloropropane	96128	VOC		X		
1,2-Dibromoethane	106934	VOC		X		
1,2-Dichlorobenzene	95501	VOC/SVOC	X			
1,3-Dichloropenzene	541731	VOC/SVOC	X			
1,4-Dichlorobenzene	106467	VOC/SVOCI	X			
3,3'-Dichloropenzidine	91941	SVOC		Х		
1,1-Dichloroethane	75343	VOC	X			
1,2-Dichloroethane	107062	VOC	X			
1,1-Dichloroethene	75354	VOC	X			
1.2-Dichloroethene (c:s-)	156592	I VOC	X			
1,2-Dichloroethene (trans-)	156605	I VOC	X		<u> </u>	
1.2-Dichloroethene (total)	156605	VOC		X		
2.4-Dichloropnenol	120832	SVOC	X			
1,2-Dichloropropane	78875	VOC	X			
1,3-Dichloropropene (cis-)	542756	VOC	X	<u> </u>		
1,3-Dichloropropene (trans-)	542756	VOC	X			
Oieldnn	60571	PEST	X		1	
Diethyl phthalate	84662	SVOC		X		
2.4-Dimethylphenol	105679			X		
Dimethylphthalate	131113			X		
Di-n-outylphthalate	84742	SVOC		<u> </u>		
Di-n-octylphthalate	117840		X			
4.6-Dinitro-2-methylpnenoi	534521			X ·		
2.4-Dinitropnenol	51285			ı X		
2.4-Dinitrotoluene	121142			X		
2,6-Dinitrotoluene	606202	SVOC		! X		
Endosulfan I	115297	PEST		X(9)		
Endosulfan II	115297			X <sub>(3)</sub>		
Endosulfan sulfate	1031078	PEST			(endosulfan)	

# SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 3 OF 4

Chamina			Basis of Value to be Used in RI Report				
Chemical	CAS Number	Chemical Fraction	Promulgated Value <sup>(1)</sup>	Calculated Value <sup>(2)</sup>	Surrogate Calculated		
Endnn	72208	I PEST I	x		Value <sup>(3)</sup>		
Endrin aidehyde	7421363	PEST	<del></del>				
		, 23,			X		
Endnn ketone	53494705	PEST			(endrin)		
	13.3.3.		}		X		
Ethylbenzene	100414	VOC	×		(endrin)		
Fluoranthene	206440		<del></del>		<del>-</del>		
Fluorene	86737		$\frac{\hat{x}}{\hat{x}}$		<del></del>		
Heptachlor	76448	PEST	<del>- x</del>	<del></del>	· · · · · · · · · · · · · · · · · · ·		
Heptachior epoxide	1024573		$\frac{\hat{x}}{\hat{x}}$				
Hexachiorobenzene	118741	SVOC	×				
Hexachlorobutadiene	87683	SVOC		×	1		
Hexachlorocyclopentagiene	77474	SVOC		<del>^</del>	<del>!</del>		
Hexachloroethane	67721	SVOC	- x				
2-Hexanone	73663715	VOC		X	<u> </u>		
ndeno(1,2,3-cd)pyrene	193395	svoc		<del></del>			
ron	7439896	INORG	(4)	(4)	<del></del>		
sophorone	78591	SVOC			(4)		
Des	7439291	INORG		Х			
Magnesium	7439954		χ ,	(6)	<u> </u>		
Manganese	7439965	INORG			(0)		
Mercury	7439976	INORG		X			
Methoxychlor	72435	INORG	X				
Methylene chlonge	75092	PEST	X		<u> </u>		
-Methylnaphthalene		VOC	X		<u> </u>		
-Methyl-2-pentanone	91576	SVOC	<del></del>	Х			
-Methylpnenoi	108101	VOC	Х		:		
-Methylphenol	95487	SVOC		X	1		
apnthalene	106445	SVOC		X			
ickel	91203	svoc	X				
Nitroaniiine	7440020	INORG	X				
Nitroaniine	88744	SVOC		X			
Nitroaniline	99092	SVOC		X			
itropenzene	100016	SVOC		X			
Nitropnenol	98953	SVOC		X			
	88755	SVOC	1		X		
Nitropnenoi	100007	61406			(4-nitropnenol		
Nitrosociphenviamine	100027	SVOC		X	l		
Nitrosodi-n-propylamine	86306	SVOC		X			
2'-Oxypis(1-chioropropane)	621647	SVOC	<u> </u>	X			
	108601	SVOC :	31	· 3)	.51		
ntachlorophenol lenanthrene	37865	SVOC	Х				
	85018	SVOC			X		
enoi	108952	svoc	Х		(naphthalene)		
tassium							
rene	129000	SVOC	<u>.                                    </u>	:01	(ċ)		

### SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 4 OF 4

<del></del>		i	Basis of Va	lue to be Used in	RI Report
Chemical	CAS Number	Chemical Fraction	Promuigated Value <sup>(1)</sup>	Calculated Value <sup>(2)</sup>	Surrogate Calculated Value <sup>(3)</sup>
Selenium	7782492	INORG	X		
Silver	7440224	INORG	X		
Sodium	7440235	INORG	(6;	, <b>6</b> )	(6)
Styrene	1 100425	VOC	X		<u> </u>
1,1,2,2-Tetrachioroethane	79345	VOC	X		
Tetrachloroethylene	127184	VOC	X		
Thallium	6533739	INORG	X		<u> </u>
Toluene	108883	VOC	X		<del> </del>
Toxaphene	8001352	PEST	X		<u> </u>
1,2,4-Trichloropenzene	120821	SVOC		X	
1,1,1-Trichloroethane	71556	i voc	X		<u> </u>
1,1,2-Trichloroethane	79005		X	<u> </u>	<u> </u>
Trichloroethylene	79016	<del></del>	X	!	<del> </del>
2,4,5-Trichlorophenol	95954	SVOC	<u> </u>	X	1
2.4.5-Trichloropnenal	88062		<u> </u>	X	1
Vanadium	7440622		X		1
Vinyi chloride	75014		X	<u> </u>	
Xylene (total)	1330207	VOC	X		<del> </del>
Zinc	7440666	INORG	X	<u></u>	

INORG Inorganic PEST Pesticide

SVOC Semivolatile organic compound VOC Volatile organic compound

- State of Connecticut Remediation Standard Regulations, Section 22a-133k (January 1996).
- Published toxicity chteria is available. Toxicity chteria from the current USEPA Region ill Risk-Gased Concentration Table (October 22, 1997) will be used to calculate a value using the methodology presented in the State guidance (January 1996).
- 3 No toxicity criteria is available. Toxicity criteria for a similarly structured chemical (noted in parentheses) will be used to calculate a value.
- 4 Region I does not advocate a quantitative evaluation of this chemical. Exposure to this chemical will be addressed in a qualitative fashion.
- No promulgated value or published toxicity criteria are available. A similarly structured chemical with published toxicity criteria could not be identified. Exposure to this chemical will be addressed in a qualitative fashion.
- 5 Chemical is an essential nutrient.
- 7 Value for chlordane is used.
- 8 Value for hexavalent chromium is used for conservative purposes.
- 9 Value for endosulfan is used.

TABLE 1

### SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 1 OF 4

<b>a.</b>		į L		alue to be Used	
Chemical	CAS Number	Chemical Fraction	Promulgated Value <sup>[1]</sup>	Calculated Value <sup>(2)</sup>	Surrogate Calculated
			İ		Value <sup>,3)</sup>
Acenaonthene	83329	SVOC	i i	X	10.00
Acenaphthylene	208968	svoc	X		
Anthracene	120127	SVOC	X		
Acetone	67641	VOC	X		
Aldrin	309002	PEST		X	
Aluminum	7429905	INORG	(4)	(4)	(4)
Antmony	7440360	INORG	X		<del></del>
Arsenic	7440382	INORG	X		
Banum	7440393	INORG	X		<del> </del>
Benzene	71432	VOC	X		<u> </u>
enz(a)anthracene	56553	SVOC	X		
Benzo(b)fluoranmene	205992	SVOC	X		
Benzork)fluoranthene	207089	svoc	X		
Benzo(g,h,i)perylene	191242	SVOC			X
			ļ		(pyrene)
Benzola)pyrene	50328	svoc	X		
Beryllium	7440417	INORG	X		
BCH (aipha-)	319846	PEST		Х	
BCH (beta-)	319857	PEST		Х	
ВСН (дела-)	319868	PEST			(alpha-8HC)
BCH (gamma-: Lindane)	58899	PEST	X		1
Bis(2-chloroethoxy)methane	111911	SVOC	.5)	,5)	(5)
is(2-chloroethy)ether	111444	SVOC	×		
is(2-ethylhexyl)phthalate	117817	SVOC	$\frac{\hat{x}}{x}$	<del></del>	<del></del>
romochioromethane	74975	VOC		<u></u>	1 X
·		.55			(bromodichiero-
romocionloromethane	75274	VOC		X	1
romoform	75252	voc i	× 1		
romomethane	74839	voc i	· i	×	<del> </del>
-Bromoonenvi-onenviether	101553	SVOC		×	
-Butanone	78933	vac	X		
utvibenzviphthalate	85687	SVOC	X		T
admium	7440439	INORG	X		i
alcium	7440702	INORG	<b>3</b> )	(5)	:51
aroazole	86748	SVOC	i	Х	i
arbon disurfide	75150	voc i		X	i
arbon tetrachionde	56235	VOC			<del></del>
hioroane (alpha-)	57749	PEST	X <sup>1</sup> 77		<del></del>
hlordane (gamma-)	57749		X <sub>1,1</sub>		<del>-                                    </del>
-Chioroaniine	106478	PEST   SVOC	<u></u>	<del></del>	1
hioropenzene	1 108907		· · · · · · · · · · · · · · · · · · ·		<u> </u>
hiorogipromomethane	<del></del>	VOC I			<del>!</del>
hloroetnane	124481	VOC	_ (		<u> </u>
hiorotorm	75003	VOC		X	1
hloromethane	57663	VOC	<u> </u>		<del></del>
MOTOTIFERIALE	74873	70C	1	Х	

# SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 2 OF 4

			Basis of V	alue to be Used in	
Chemical	CAS	Chemical	Promuigated	Calculated	Surrogate
	Number .	Fraction	Value.1)	∀alue <sup>(2)</sup>	Calcutated Value <sup>(3)</sup>
2-Chloronaonthalene	91587	SVOC		X	
2-Chloroph <b>enol</b>	95578	SVOC	Х		
4-Chlorophenyi-pnenyiether	7005723	svoc			X (4-Bromophenyl- phenylether)
Chromium (total)		INORG	X <sup>(8)</sup>		
Chrysene	218019	svoc		X	
Cobalt	7440484	INORG		X	
Copper	7440508	INORG	(4)	(4)	(4)
4.4'-ODD	72548	PEST		X	
4,4'-DDE	72559	PEST		Х	
4.4'-DDT	50293	PEST		Х	
Dibenzoturan	132649	SVOC		Х	
Dibenz(a,h)anthracene	53703	SVOC		Х	
1.2-Dibromo-3-chloroprocane	96128	voc		X	1
1.2-Dibromoethane	106934	voc		X	
1,2-Dichlorobenzene	95501	VOC/SVOC	X		
1,3-Dichlorobenzene	541731	voc/svoc	X		<u> </u>
1.4-Dichlorobenzene	106467	voc/svoc	Х		
3.3'-Dichlorobenziaine	91941	SVOC		X	1
1,1-Dichloroethane	75343	VOC	Х		
1.2-Dichloroethane	107062	VOC	X		
1,1-Dichlorcetnene	75 <b>354</b>	VOC	X		
1.2-Dichloroethene (cis-)	156592	VOC	X		
1 2-Dichloroethene (trans-)	156605	VOC	X		1
1 2-Dichloroethene (total)	156605			X	<u> </u>
2.4-Dichiaroanenai	120832	SVOC	<u> </u>		
1.2-Dichioropropane	78875			<u> </u>	<u> </u>
1 3-Dichlorogropene (cis-)	542756	VOC	X	<u> </u>	
1,3-Dichloropropene (trans-)	542756	VOC	Ж		
Cieldnn	60571	PEST	X	<u> </u>	
Diethyl chthalate	84662	SVOC		X	
2,4-Dimethylphenol	105679	SVOC		<u> </u>	
Dimethylphthalate	131113	SVOC		X	
Di-n-but/tonthalate	84742	SVOC	X		<del>-  </del>
Di-n-octylonthalate	117840	SVOC	Х		<del> </del>
4 3-Dinitro-2-methylphenai	534521	SVOC		<u> </u>	
2,4-Dinitrophenoi	51285			i X	<del></del>
2.4-Dintrotoluene	121142				-
2,6-Dintrotoluene	606202			X	1
Endosulfan I	115297	PEST		X <sub>(8)</sub>	<u> </u>
Endosurfan II	115297	PEST		X <sub>(8)</sub>	_!
Endosufan suffate	1031078	PEST			(endosurfan)
ndnn	72208	PEST	Y	1	
Endrin aldenyde	7421363	PEST			(endnn)
narin ketone	53494705	PEST			X engnni

TABLE 1

### SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 3 OF 4

		i i	Basis of V	Basis of Value to be Used in RI Report				
Chemical	CAS	Chemical	Promulgated	Calculated	Surrogate			
	Number	Fraction	Value <sup>(1)</sup>	Value <sup>(2)</sup>	Calculated Value <sup>(3)</sup>			
Ethylbenzene	100414	voc	×		1			
Fluoranthene	206440	SVOC	X I					
Fluorene	86737	SVOC	X					
Heotachior	76448	PEST	Х					
Heptachior epoxide	1024573	PEST	X					
Hexachiorobenzene	118741	SVOC	X					
Hexachiorobutadiene	87683	svoc		X				
Hexachiorocyclopentadiene	77474	SVOC		X				
Hexachioroethane	67721	SVOC	X 1					
2-Hexanone	73663715	voc		X				
indenoi1.2.3-cd)byrene	193395		i	X				
Iron	7439896	INORG	41	4)	(4)			
Isophorone	78591			×	<del> </del>			
Lead	7439291	INORG	X.	· · · · · · · · · · · · · · · · · · ·	<del> </del>			
Magnesium	7439954	INORG	- <u>(</u>	(6)	(6)			
Manganese	7439954	INORG		×	<del> </del>			
Mercury	7439905	INORG	<del></del>	^				
Methoxycniar	72435				<u> </u>			
Methylene chlonde	75092	PEST	<del></del>		<del> </del> _			
2-Methynaphthalene	91576	SVOC	<del></del>	X	<del> </del>			
4-Methyl-2-pentanone	108101	VOC	x	<del></del>	<del> </del>			
2-Methylphenol	95487	SVOC		X	<del> </del>			
4-Methyphenol	106445			<del></del>	<u> </u>			
Naphthalene		SVOC	<del>x</del>	^				
Nickel	7440020	SVOC	<del></del>		<u> </u>			
2-Nitroaniine	88744	SVOC	<del></del>	×	<del></del>			
3-Naroaniine			1	<del></del>				
≟-Nitroaniine	99092	SVOC			!			
Nitropenzene	1 00016	SVOC	į	<u> </u>	1			
2-Nitrochenoi	98953	SVOC		X	1			
2-1410 Chenoi	88755	svoc			X (4 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2			
4-Nitrochenol	100027			<del></del>	[4-nitrophenoi)			
N-Nitrosogiphenylamine	1 100027 1	SVOC		<u>X</u>	!			
N-Nitrosogi-n-propylamine	86306	SVOC		X				
	521647	svoc	5)	X	<u>!</u>   5			
2.2'-Oxycis(1-chlorocropane)	108601	svoc	!	<del></del>	:31			
Pentachiorophenoi	87865	SVOC	X		<u> </u>			
Phenanthrene	35018	SVOC	<u> </u>		1			
Phenoi	108952	SVOC	χ 1		i			
Potassium	7440097	:NORG	ŧ,	<b>:</b>	, d)			
<sup>2</sup> vrene	1 129000 1	SVOC	τ,		i			
Saleniim	7732492	NORG						
Silver	7440224	NORG I	4 ;		1			
Sodium	7440235	NORG	\$i	έį	;Đ)			
Styrene	100425	VOC	<u> </u>					
1,1,2,2-Tetrachioroethane	79345	VOC	* 1		<u> </u>			
Tetrachioroethylene	127184	voc	<del>``</del>		<del>i</del>			
hallium	6533739	NORG	<del>-</del> X		1			
			<del></del>		<del></del>			
foluene	1088831	.cc	•		i contract of the contract of			

### SOURCE OF CONNECTICUT REMEDIATION STANDARDS CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 4 OF 4

			Basis of V	alue to be Used in	RI Report
Chemical	CAS Number	Chemical Fraction	Promuigated Value <sup>(1)</sup>	Calculated Value <sup>(2)</sup>	Surragate Calculated Value <sup>(3)</sup>
1.2.4-Trichlorobenzene	120821	SVOC		X	
1,1,1-Trichloroethane	715 <b>56</b>	voc	X		
1,1,2-Trichloroethane	79005	voc	Χ		
Trichloroethylene	79016	voc	Х		
2.4.5-Trichlorophenol	95 <b>954</b>	SVOC		X	
2,4,6-Trichlorophenol	88062	SVOC		X	
Vanadium	7440622	INORG	Х		
Vinyl chloride	75014	voc	Х		
Xylene (total)	1330207	voc	X		
Zinc	7440666	INORG	X		

INORG Inorganic PEST Pesticide

SVOC Semivolatile organic compound VOC Volatile organic compound

- 1 State of Connecticut Remediation Standard Regulations, Section 22a-133k (January 1996).
- Published toxicity criteria is available. Toxicity criteria from the current USEPA Region III Risk-Based Concentration Table (October 22, 1997) will be used to calculate a value using the methodology presented in the State guidance (January 1996).
- 3 No toxicity criteria is available. Toxicity criteria for a similarly structured chemical (noted in parentheses) will be used to calculate a value.
- 4 Region I does not advocate a quantitative evaluation of this chemical. Exposure to this chemical will be addressed in a qualitative fashion.
- 5 No promulgated value or published toxicity criteria are available. A similarly structured chemical with published toxicity criteria could not be identified. Exposure to this chemical will be addressed in a qualitative fashion.
- 5 Chemical is an essential nutrient.
- Value for chlordane is used.
- 8 Value for hexavalent chromium is used for conservative purposes.
- 9 Value for endosulfan is used.

TABLE 2

### CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 1 OF 3

	Published Toxclol	ogical Criteria <sup>(1)</sup>	Calculated Remediation Standards <sup>(2)</sup>					
Chemical	RfD <sub>oral</sub>	CSF <sub>utel</sub>			ng/kg)		Groundwater (ug/L)	
	(mg/kg/day)	(kg/day/mg)	RES DE(3)	NC DE(1)	GA/GAA PM	GB PM	GA/GAA	
	'''	\ <b>.</b>					GP	
Acenaphthene	6.00E-02	NA NA	1000 <sup>(4)</sup>	2500(4)	8.4	84	420	
Aldrin	3.00E-05	1 70E+01	0.036	0.34	0.000041	0.00041	0.0021	
Benzo(g,h,i)perylene	NA	NA	1000 <sup>(5)</sup>	2500 <sup>(5)</sup>	5.6 <sup>(5)</sup>	56 <sup>(5)</sup>	280(5)	
BCH (alpha-)	NA	6 30E+00	0.097	0.91	0.00011	0.0011	0.0056	
BCH (beta-)	NA	1 80E+00	0.34	3.2	0.00039	0.0039	0.0194	
BCH (delta-)	NA	NA	0 097(6)	0.91(6)	0.00011(8)	0.0011(6)	0.0056 <sup>(6)</sup>	
Bromochloromethane	NA	NA	47 <sup>(7)</sup>	440 <sup>(7)</sup>	0.054 <sup>(7)</sup>	0.54 <sup>(7)</sup>	2 7(7)	
Bromodichloromethane	2.00E-02	6 20E-02	9.9	92	0.011	0.11	0 56	
Bronomethane	1.40E-03	NA	95	1000(4)	0.2	2	9.8	
4-Bromophenyl-phenylether	5.80E-02	NA	500 <sup>(4)</sup>	1000(4)	8.2	82	410	
Carbazole	NA	2 00E-02	31	290	0.036	0.36	1,8	
Carbon disulfide	1.00E-01	NA	500 <sup>(4)</sup>	1000 <sup>(4)</sup>	14	140	700	
4-Chloroaniline	4.00E-03	NA	270	2500(4)	0.56	5.6	28	
Chloroethane	4.00E-01	2 90E-03	210	1000(4)	0.24	2 4	12	
Chloromethane	NA	1 30E-02	47	440	0.054	0 54	27	
4-Chloro-3-methylphenoi	NA NA	NA	1000(8)	2500 <sup>(0)</sup>	7(6)	70 <sup>(6)</sup>	350 <sup>(8)</sup>	
2-Chloronaphthalene	8 00E-02	NA NA	1000(4)	2500(4)	11	110	560	
4-Chlorophenyl-phenylether	NA NA	NA NA	500 <sup>(9)</sup>	1000(9)	8.2(9)	82 <sup>(6)</sup>	410(9)	
Chrysene	NA NA	7 30E-03	84	780	0.096	0.96	4.8	
Cobalt	6.00E-02	NA NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	2200(16)(11)	22000 <sup>(10)(11)</sup>	420	
4.4'-DDD	NA NA	2 40E-01	2.6	24	0.0029	0.029	0 15	
4.4' DDE	NA NA	3 40E-01	1.8	17	0 0021	0.021	01	
4.4' DDT	5.00E-04	3 40E-01	1.8	17	0 0021	0 021	01	
Dibenzofuran	4.00E-03	NA	270	2500(4)	0 56	56	28	
Dibenz(a,h)anthracene	NA	7 30E+00	0.084	0.78	0 000096	0 00096	0 0048	
1.2-Dibromo-3-chloropropane	NA NA	1.40E+00	0.44	4.1	0.0005	0 005	0 025	
1 2-Dibromoethane	NA	8 50E+01	0 0072	0.067	0 0000082	0.000082	0 00041	
3 3'-Dichlorobenzidine	- NA	4 50E-01	1.4	13	0.0016	0 0 1 6	0 078	
1.2 Dichloroethene (total)	2.00E-02	NA	500 <sup>(4)</sup>	1000(4)	2 8	28	140	
Diethyl phthalate	8 00E·01	NA	1000(4)	2500(4)	110	1100	5600	

TABLE 2

## CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 2 OF 3

	Published Toxclo	logical Criteria <sup>(1)</sup>					
Chemical	RID	CSF <sub>oral</sub>			Groundwater (ug/L)		
	(mg/kg/day)	(kg/day/mg)	RES DE(1)	NC DE(3)	GA/GAA PM	GB PM	GA/GAA
2.1.0							GP
2,4-Dimethylphenol	2 00E-02	NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	2,8	28	140
Dimethylphthalate	1.00E+01	NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	1400	1400	70000
4.6-Dinitro-2-methylphenol	1 00E-04	NA	6.8	200	0 014	0 14	0.7
2,4-Dinitrophenol	2 00E-03	NA	140	2500 <sup>(4)</sup>	0.28	28	14
2,4-Dinitrotoluene	2 00E-03	NA	140	2500 <sup>(4)</sup>	0.28	2.8	14
2,6-Dinitrotoluene	1.00E-03	NA	68	2000	0.14	1.4	7
Endosulfan j	6.00E-03	NA	410	1200	0.84	8.4	42
Endosulfan II	6.00E-03	NA	410	1200	0.84	8.4	42
Endosulfan sulfate	NA	NA	410 <sup>(12)</sup>	1200(12)	0.84(12)	8.4(12)	42(12)
Endrin aldehyde	NA	NA	20(13)	610 <sup>(13)</sup>	NE(13)	NE <sup>(13)</sup>	NE <sup>(13)</sup>
Endrin ketone	NA	NA	20(13)	610 <sup>(13)</sup>	NE <sup>(13)</sup>	NE <sup>(13)</sup>	NE(13)
dexachlorobutadiene	2.00E-04	7.80E-02	7.9	73	0.009	0.09	0.45
lexachlorocyclopentadiene	7.00E-03	NA	470	2500(4)	0.98	9.8	49
2-Hexanone	4.00E-02	NA	500 <sup>(4)</sup>	1000(4)	5.6	56	280
Indeno(1,2,3-cd)pyrene	NA	7 30E 01	0.84	7.8	0.00096	0.0096	0.045
Isophorone	2 00E-01	9 50E-04	640	2500(4)	0.74	7.4	37
Manganese	2.30E-02	NA	1600	47000	50(10)(14)	500 <sup>(10)(14)</sup>	160
2-Methylnaphthalene	4.00E-02	NA NA	1000 <sup>(4)</sup>	2500(4)	5.6	56	280
2-Methylphenot	5 00E-02	NA	1000(4)	2500(4)	7	70	350
4-Methylphenol	5.00E-03	NA NA	340	2500 <sup>(4)</sup>	0.7	7	35
2-Nitroaniline	6 ONE-05	NA	4.1	1200	0.0084	0.084	0.42
3-Nitroaniline	3 00€-03	NA	200	2500 <sup>(4)</sup>	0 42	4.2	21
-Nitroaniline	3 00E-03	NA -	200	2500 <sup>(4)</sup>	0 42	4.2	21
litrobenzene	5 00E-04	NA -	34	1000	0.07	0.7	3.5
Nitrophenol	NA NA	NA NA	540(15)	2500(15)	1.1(13)	11(15)	56(15)
-Nitrophenol	8 00E-03	NA NA	540	2500 <sup>(4)</sup>	11	11	56
Nitrosodiphenylamin	NA	4 90E-03	130	1200	0.14	14	71
l-Nitrosodi-n-propylamma	NA NA	7.00E+00	0.088	0.82	0 0001	0 00 i	0.005
Phenanthrene	NA	NA NA	1000 <sup>(5)</sup>	2500 <sup>(5)</sup>	5 6 <sup>(5)</sup>	56 <sup>(5)</sup>	280(5)
1,2,4 Trichlorobenzene	1 00E 02	NA NA	680	2500 <sup>(4)</sup>	14	14	70

1

### CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 3 OF 3

	Published Toxclo	logical Criteria <sup>(1)</sup>	Calculated Remediation Standards <sup>(2)</sup>					
Chemical	RfD <sub>oral</sub>	CSF <sub>urai</sub>		Groundwater (ug/L)				
	(mg/kg/day)	(kg/day/mg)	RES DE <sup>(3)</sup>	AC DE(a)	GA/GAA PM	A PM GB PM		
2,4,5-Trichlorophenol	1.00E-01	NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	14	140	700	
2,4,6-Trichlorophenol	NA	1.10E-02	56	520	0.064	0 64	3.2	

RID

Reference dose

CSF

Cancer slope factor

**RES DE** 

Direct exposure criteria for residential land use

I/C DE

Direct exposure criteria for industrial/commercial land use.

GA/GAA PM

Pollulant mobility criteria for a GA/GAA classified area

GB PM

Pollutant mobility criteria for a GB classified area

GA/GAA GP

Groundwater protection criteria for a GA/GAA classified area

NA

Not available

NE

None established by Connecticut DEP (January 1996)

- 1 Values obtained from current USEPA Region III Risk-Based Concentration Table (October 22, 1997)
- 2 Calculated using methodologies presented in State guidance (January 1996).
- 3 Calculated value for direct exposure for volatile and semivolatile organics is replaced with the appropriate ceiling limit if the calculated value exceeds the ceiling limit. Ceiling limit for volatiles is 500 mg/kg for residential exposure and 1000 mg/kg for industrial/commercial exposure. Ceiling limit for semivolatiles is 1000 mg/kg for residential exposure and 2500 mg/kg for industrial/commercial exposure
- 4 Ceiling limit. Calculated value exceeds the ceiling limit.
- 5 Value for naphthalene is used
- 6 Value for alpha-BHC is used
- 7 Value for chloromethane is used.
- 8 Value for 3-methylphenol is used.
- 9 Value for 4-bromophenyl-phenylether is used.
- 10 Value is for aqueous units (ug/l.) and is based on SPLP or TCLP analytical results.
- 11 Value is based on the Region III RBC for tap water (2200 ug/L).
- 12 Value for endosulfan is used
- 13 Value for endrin is used.
- 14 Value is based on the secondary Federal MCL for drinking water (50 ug/L).
- 15 Value for 4-nitrophenol is used.

TABLE 2

## CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 1 OF 3

	Published Toxclo	ogical Criteria <sup>(1)</sup>	Calculated Remediation Standards <sup>(3)</sup>						
Chemical	RfD <sub>ural</sub> (mg/kg/day)	CSF <sub>ore</sub> , (kg/day/mg)		Soll (mg/kg)					
	(gg/cay)	(kg/day/mg)	RES DE <sup>(3)</sup>	NC DE(3)	GA/GAA PM	GB PM	(ug/L) GA/GAA		
Acenaphthene	6 00E-02	NA NA	1000 <sup>(4)</sup>		-		GP		
Aldrin	3.00E-05	1 70E +01	0.036	2500 <sup>(4)</sup>	8.4	84	420		
Benzo(g.h.i)perylene	NA	NA NA	1000(5)	0.34	0.000041	0.00041	0 0021		
BCH (alpha-)	NA -	6 30E +00	0.097	2500 <sup>(5)</sup>	4(5)	40 <sup>(5)</sup>	200 <sup>(5)</sup>		
BCH (beta-)	NA NA	1.80E+00	0.34	0.91	0.00011	0 0011	0.0056		
BCH (delta-)	NA	NA NA	0.097 <sup>(6)</sup>	3.2	0.00039	0.0039	0.0194		
Bromochloromethane	NA			0.91(6)	0.00011 <sup>(6)</sup>	0.0011(6)	0 0056 <sup>(6)</sup>		
Bromodichloromethane	2 00E-02	NA NA	9.9 <sup>(7)</sup>	92(1)	0.011(7)	0.11(7)	0.56(7)		
Bromomethane		6 20E-02	9.9	92	0.011	0.11	0.56		
4 Bromophenyl phenylether	1.40E-03	NA NA	95	1000(4)	0.2	2	9.6		
Carbazole	5 80E-02	NA NA	500(4)	1000(4)	8 2	82	410		
arbon disulfide	NA	2.00E-02	31	290	0.036	0.36	1.8		
1 Chloroanitine	1 00E-01	NA	500 <sup>(4)</sup>	1000(4)	14	140	700		
Chloroethane	4.00E-03	NA	270	2500(4)	0.56	5 6	28		
hloromethane	4 00E-01	2.90E-03	210	1000(4)	0.24	24			
	- NANA	1 30E-02	47	440	0.054	0.54	12 7		
Chloro-3-methylphenal	NA NA	NA NA	NA <sup>(a)</sup>	NA <sup>(b)</sup>	NA <sup>(6)</sup>	NA <sup>(6)</sup>	NA <sup>(8)</sup>		
2 Chloronaphthalene	8 00E-02	AII	1000(4)	2500[1]	11				
Chlorophenyl phenylether	NA	AI1	500 <sup>(9)</sup>	1000(9)	8 2(9)	110	560		
tuysene	NA NA	7 30E-03	84	780	·   ——— -	82(9)	410 <sup>(9)</sup>		
Coball	6 00E-02	NA NA	1000(4)	2500(4)	0.096	0.96	48		
4DDD	NA -	2 40E-01	2.6	250,7	420(10)	4200(10)	420		
4'-DDE	NA NA	3 40E-01	1.8	17	0 0029	0.029	0.15		
4'-DDT	5.00E-04	3.40E-01	1.8	l ——— :	0.0021	0.021	01		
ibenzofuran	4 00E-03	NA NA	270	17	0.0021	0.021	0.1		
ibenz(a,h)anthracene	NA NA	7 30E +00		2500 <sup>(4)</sup>	0 56	5 6	28		
2-Dibromo-3 chloropropane	NA -	1 40E+00	0.084	0 78	0.000096	0 00096	0.0048		
2-Dibromosthane	NA -	8 50E+01	0 44	4 1	0 0005	0 005	0.025		
3' Dichlorobenzidine		4 50E-01	0 0072	0 067	0.0000082	0 000082	0.00041		
2 Dichloroethone (total)	9.00E-03		1 4	13	0.0016	0.016	0.078		
ethyl phthalate		NA	500(4)	1000(4)	12	12	63		
4-Dimethylphenol	8 00E-01	NA NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	110	1100	5600		
wriothylphthalato	2 00E-02	NA NA	1000(4)	2500[4]	2,8	28	140		
and the mark	1 00E+01	NA	1000(4)	2500(4)	1400	14000	70000		

TABLE 2

#### CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 2 OF 3

	Published Toxcloi	opical Criteria(1)	Calculated Remediation Standards <sup>(2)</sup>						
					Groundwater (ug/L)				
Chemical	RID	CSF			ng/kg)	GB PM	GA/GAA		
	(mg/kg/day)	(kg/day/mg)	RES DE <sup>(3)</sup>	NC DE(3)	GA/GAA PM	GB PM	GP		
					0.014	0.14	07		
,6-Dinitro-2-methylphenol	1.00E-04	NA	6.8	200		2.8	14		
,4-Dinitrophenol	2.00E-03	NA NA	140	2500(4)	0.28		14		
4-Dinitrotoluene	2.00E-03	NA NA	140	2500 <sup>(4)</sup>	0.28	2.8	7		
2,6-Dinitrotoluene	1.00E-03	NA	68	2000	0.14 0.64	B.4	42		
ndosulfan l	6 00E-03	NA	410	1200	0.84	8.4	42		
ndosulfan li	6 00E-03	NA	410	1200	0.84(12)	8.4(12)	42(12)		
ndosulfan sulfate	NA	NA	410 <sup>(12)</sup>	1200(12)	NE <sup>(13)</sup>	NE <sup>(13)</sup>	NE <sup>(13)</sup>		
Endrin aldehyde	NA	NA	20(13)	610(13)	NE'	NE(13)	NE(13)		
Endrin ketone	NA	NA	20(13)	610 <sup>(13)</sup>	NE <sup>(13)</sup>	0.09	0.45		
lexachlorobutadiene	2.00E-04	7 80E-02	7.9	73	0.009		49		
lexachlorocyclopentadiene	7.00E-03	NA	470	2500 <sup>(4)</sup>	0.98	9.8	280		
2-Hexanone	4.00E-02	NA	500 <sup>(4)</sup>	1000(4)	5.6	56 0.0096	0 045		
ndeno(1,2,3-cd)pyrene	NA	7.30E-01	0.84	7.8	0.00096	7.4	37		
sophorone	2 00E-01	9.50E-04	640	2500 <sup>(4)</sup>	0.74 50 <sup>(10</sup> (14)	500 <sup>(10)(14)</sup>	160		
Manganese	2.30E-02	NA	1600	47000			280		
2-Methylnaphthalene	4.00E-02	NA	1000(4)	2500 <sup>(4)</sup>	5.6	56	350		
2-Methylphenol	5 00E-02	NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	7	70	35		
4-Methylphenol	5.00E-03	NA	340	2500 <sup>(4)</sup>	0.7	7	0.42		
2-Nitroaniline	6.00E-05	NA	4.1	1200	0.0084	0.084	21		
3-Nitroaniline	3.00E-03	NA	200	2500 <sup>[4]</sup>	0.42	4.2			
	3.00E-03	NA	200	2500 <sup>[4]</sup>	0.42	4.2	3.5		
4-Nitroaniline	5.00E-04	NA NA	34	1000	0.07	0.7	56 <sup>(15)</sup>		
Nitrobenzene	NA	NA NA	540(15)	2500(15)	1.1 <sup>(15)</sup>	11(15)			
2-Nitrophenol	1	NA NA	540	2500 <sup>(4)</sup>	1.1	11	56		
4-Nitrophenol	8.00E-03 NA	4.90E-03	130	1200	0.14	1.4	7.1		
N-Nitrosodiphenylamine	NA	7.00E+00	0 088	0.82	0.0001	0.001	0 005		
N-Nitrosodi-n-propylamine	1.00E-02		680	2500 <sup>(4)</sup>	1.4	14	70		
1,2,4-Trichlorobenzene		NA NA	1000 <sup>(4)</sup>	2500 <sup>(4)</sup>	14	140	700		
2.4.5-Trichlorophy	1.00E-01 NA		56	520	0 064	0 64	32		

Reference dose RID

Cancer slope factor CSF

#### CALCULATED AND SURROGATE CALCULATED VALUES CTO 260 LOWER SUBASE RI NEW LONDON, GROTON, CONNECTICUT PAGE 3 OF 3

RES DE

Direct exposure criteria for residential land use

I/C DE

Direct exposure criteria for industrial/commercial land use.

**GA/GAA PM** 

Pollutant mobility criteria for a GA/GAA classified area

GB PM

Pollutant mobility criteria for a GB classified area

**GA/GAA GP** 

Groundwater protection criteria for a GA/GAA classified area

NA Not available

NE

None established by Connecticut DEP (January 1996)

- 1 Values obtained from current USEPA Region III Risk-Based Concentration Table (October 22, 1997)
- 2 Calculated using methodologies presented in State guidance (January 1996).
- 3 Calculated value for direct exposure for volatile and semivolatile organics is replaced with the appropriate ceiling limit if the calculated value exceeds the ceiling limit. Ceiling limit for volatiles is 500 mg/kg for residential exposure and 1000 mg/kg for industrial/commercial exposure. Ceiling limit for semivolatiles is 1000 mg/kg for residential exposure and 2500 mg/kg for industrial/commercial exposure.
- 4 Ceiling limit. Calculated value exceeds the ceiling limit.
- 5 Value for pyrene is used.
- 6 Value for alpha-BHC is used.
- 7 Value for bromodichloromethane is used.
- 8 Chemical will be addressed qualitatively at CTEP's request
- 9 Value for 4-bromophenyl-phenylether is used.
- 10 Value is for aqueous units (ug/L) and is based on SPLP or TCLP analytical results.
- 11 Value is based on the Region III RBC for tap water (2200 ug/L).
- 12 Value for endosultan is used.
- 13 Value for endrin is used.
- 14 Value is based on the secondary Federal MCL for drinking water (50 ug/L).
- 15 Value for 4-nitrophenol is used.



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C-49-03-8-156

March 20, 1998

Brown & Root Environmental Project Number 7237

Mr. Mark Lewis
Connecticut Department of Environmental Protection
Water Management Bureau
Permitting, Enforcement, and Remediation Division
Federal Remediation Program
79 Elm Street
Hartford, Connecticut 06106-5127

Reference:

CLEAN Contract No. N62472-90-D-1298

Contract Task Order No. 0260

Subject:

Responses to CTDEP's Comments on Calculated Remediation Standards

Lower Subase Remedial Investigation

Naval Submanne Base - New London, Groton, Connecticut

Dear Mr. Lewis:

Brown & Root (B&R) Environmental and the Navy received your February 27, 1998 comment letter regarding the Remediation Standards that were calculated for use in the Lower Subase Remedial Investigation. Responses to CTDEP's comments have been prepared and the appropriate revisions have been made to Tables 1 and 2, which were previously enclosed in B&R Environmental's December 23, 1997 letter. B&R Environmental, on the behalf of the United States Navy, Northern Division Facilities Engineering Command and Naval Submarine Base - New London, has enclosed the Navy's responses to CTDEP's comments and the revised tables for your review and approval.

If you have any questions regarding the responses or the information provided in the revised tables, please contact Mr. Mark Evans at (610) 595-0567 (ext. 162) or me at (412) 921-8244. It is anticipated that any remaining issues can be resolved during a conference call.

Very truly yours.

Cory AL RIGHT P.E.

Project Manager

Enclosure(s)

: Mr. Roger Boucher, NORTHDIV (letter only)

Mr. Mark Evans, NORTHDIV

Mr. Andy Stackpole, NSB-NLON Environmental

Mr. John Trepanowski, B&R Environmental

Mr. Daryl Hutson, 8&R Environmental (letter only)

Ms. Karen Smecker, B&R Environmental

File: CTO 0260

# RESPONSES TO CTDEP'S COMMENTS (2/27/98) ON THE CALCULATED CTDEP REMEDIATION STANDARDS (12/23/97) CTO 260 - LOWER SUBASE REMEDIAL INVESTIGATION NAVAL SUBMARINE BASE-NEW LONDON, GROTON, CONNECTICUT MARCH 20, 1998

### I. SURROGATE CHEMICALS USED TO SUPPLY TOXICITY VALUES

#### Comment:

1. The Navy has used naphthalene as a surrogate to represent the toxicity of benzo(g,h,l)perylene. As noted in Dr. Ginsberg's memorandum, pyrene (RfD 0.03 mg/kg/d) is a more appropriate surrogate. The RfD for naphthalene has been withdrawn from IRIS. Please recalculate the direct exposure, pollutant mobility, and ground water protection criteria for benzo(g,h,i)perylene using this approach. This approach is appropriate for a screening level risk assessment. However, the uncertainties involved with this approach should be acknowledged if these two chemicals are found to be major risk drivers at the site.

#### Response:

The direct exposure, pollutant mobility, and groundwater protection criteria for benzo(g,h,i)perylene will be recalculated using pyrene as a surrogate. Benzo(g,h,i)perylene was detected in soil and groundwater at the Lower Subase but was not found to be a major risk driver at any of the zones that were evaluated in the risk assessment. Benzo(g,h,i)perylene was only identified as a COC in groundwater at Zone 4 where it was detected in one sample at a concentration exceeding the State's Ambient Water Quality Criteria (AWQC) for the protection of human health. Consequently, this does not have any impact on the human health risk assessment.

#### Comment:

2. It is unclear why the Navy calculated criteria for phenanthrene since the regulations list direct exposure, pollutant mobility, and groundwater protection criteria for this compound. Please use the criteria listed in the Regulations for this compound. The Navy should either withdraw their request for approval of criteria for phenanthrene, or, if the Navy is requesting approval of alternative criteria for this compound under the Regulations, the Navy should so state.

#### Response:

The Navy retracts its request for approval of criteria for phenanthrene. The promulgated criteria for phenanthrene were used in the selection of COCs in the human health risk assessment. Consequently, this does not have any impact on the human health risk assessment.

#### Comment:

3. Bromodichloromethane should be used as a surrogate for bromochloromethane. Please use the criteria calculated for bromodichloromethane in place of those suits susing chloromethane as a surrogate.

#### Response:

Bromodichloromethane will be used as a surrogate for mochloromethane. Bromodichloromethane was not detected in soil and groundwater samples or any of the zones evaluated in the human health risk assessment, consequently this does not have any impact on the analysis.

#### Comment:

4. The Navy's proposal to use 3-methylphenol as a surrogate for 4-chloro-3-methylphenol is not appropriate, due to structural differences between the two compounds. The use of a qualitative risk assessment would be acceptable assuming that concentrations of this chemical dp not exceed the low part-per-billion range. Please see Dr. Ginsberg's comments for additional details.

#### Response:

No criteria will be developed for 4-chloro-3-methylphenol. Instead, as suggested, 4-chloro-3-methylphenol will be evaluated qualitatively. 4-Chloro-3-methylphenol was only detected in one soil sample at the Lower Subase and at a low concentration (34 ppb), consequently, this does not have any impact on the human health risk assessment.

### II. INCORRECT OR UNSUPPORTED POTENCY VALUES

#### Comment:

5. Several of the CSFs or RfDs used by the Navy appeared to be incorrect, based on a comparison to the values listed in the EPA Region III Risk Based Concentrations table. IRIS, or HEAST. Please recalculate the direct exposure, pollutant mobility, and ground water protection criteria using correct values for total 1,2-dichloroethene. Please assume that this value pertains to the mixture of cis and trans isomers. The RfD for the mixture should be 9E-3 mg/kg/d.

#### Response:

The direct exposure, pollutant mobility, and groundwater protection criteria for total 1.2-dichloroethene will be recalculated using an oral reference dose of 9E-3 mg/kg/day. This revision does not impact the human health risk assessment since all detected concentrations of total 1.2-dichloroethene are less than the recalculated criteria.

#### Comment:

The Department was unable to verify the potency factors listed by the Navy for several chemicals. Please either provide references to support the listed potency factors, or derive criteria using acceptable surrogates for the following compounds: chloroethane, 4,6-dinitro-2-methylphenol, 2-hexanone, and 2-methylnaphthalene. Please note that naphthalene is not an appropriate surrogate for 2-methylnaphthalene as the RfD for naphthalene has been withdrawn from IRIS. Please refer to Dr. Ginsberg's memo for additional guidance.

#### Response:

The toxicity criteria for chloroethane, 4,6-dinitro-2-methylphenol, 2-hexanone, and 2-methylphenol were obtained from the current U.S. EPA Region III Risk-based Concentration (RBC) Table dated October 22, 1997. The RBC table cites EPA's National Center for Environmental Assessment (NCEA) as the source for the values for chloroethane, 4,6-dinitro-2-methylphenol, and 2-methylphenol. Although not cited in the RBC table, EPA Region III stated in telephone call on March 12, 1998, that NCEA is also the source for the toxicity criteria for 2-hexanone. Therefore, there are no changes necessary to the proposed values.

#### Comment:

7. The Department was unable to verify the RfD listed by the Navy for 4-nitrophenol (8.00E-3 mg/kg/d). Please either provide a reference for the listed value, or use the default RfD currently listed in the RBC tables (6.2E-2 mg/kg/d).

#### Response:

The current RBC table lists 8.00E-3 mg/kg/day as the oral RfD for 4-nitrophenol and cites EPA's NCEA as the source for the value. The value of 6.2E-2 mg/kg/day was listed in the previous, outdated version of the RBC table. Therefore, there are no changes necessary to the proposed criteria.

#### III. POLLUTANT MOBILITY CRITERIA FOR METALS

#### Comment:

8. The ground water protection criterion for cobalt was calculated correctly by the Navy. However, the approach used by the Navy in calculating pollutant mobility criteria for cobalt is unacceptable. Rather than using the calculated ground water protection criterion (420 µg/l) to establish a pollutant mobility criterion for cobalt, the Navy used the EPA Region III Risk Based Criteria for tap water (2,200 µg/L) as the GAA/GA pollutant mobility criterion. This approach is less conservative than using the calculated ground water protection criterion. The correct pollutant mobility criteria for cobalt, based on the groundwater protection criteria calculated by the Navy, are 420 µg/L for a GAA/GA area, and 4,200 µg/L for a GB area (measurement by TCLP or SPLP).

#### Response:

The pollutant mobility criteria for cobalt will be changed to 420  $\mu$ g/L for a GAA/GA area and 4,200  $\mu$ g/L for a GB area. This revision has no impact on the human health risk assessment because of the following reasons: (1) none of the historical soil samples that were analyzed by TCLP had leachates that were analyzed for cobalt, and (2) only the soil samples from Zone 6 had SPLP leachates that were analyzed for cobalt and all of the results were nondetects.

#### Comment:

9. The ground water protection criterion for manganese was calculated correctly by the Navy. Rather than using the calculated ground water protection criterion (160 µg/l) to establish a pollutant mobility criterion for manganese, the Navy used the EPA Secondary MCL for drinking water (50 µg/L) as the GAA/GA pollutant mobility criterion. This approach is acceptable as it is more conservative than using the calculated ground water protection criterion.

#### Response:

No response required.

### IV. GB POLLUTANT MOBILITY CRITERIA FOR DIMETHYLPHTHALATE

#### Comment:

10. The GB pollutant mobility criteria listed for dimethylphthalate (1,400 mg/kg) in the Navy's Table 2 appears to be a typo. The correct value should be listed as 14,000 mg/kg.

#### Response:

The GB pollutant mobility criteria for dimethylphthalate will be corrected to 14,000 mg/kg. This revision has no impact on the analysis since dimethylphthalate was not detected in soil samples in any of the zones that were evaluated in the human health risk assessment.

#### V. BIS(2-CHLOROETHOXY)METHANE

#### Comment:

11. The Navy proposes a qualitative risk assessment for this compound. This approach is acceptable provided that the compound is not present at concentrations above the low part-per-billion range. As noted by Dr. Ginsberg, if it is present above this range, a more quantitative risk assessment may be required.

#### Response:

Bis(2-chloroethoxy) methane was not detected in soil or groundwater samples for any of the zones evaluated in the human health risk assessment, consequently this does not have any impact on the analysis.

Appendix F (Continued):
Human Health Risk Assessment Supporting Documentation
Sections: F.3 through F.7

(pages 90-218)

are available
in a separate file (size: 5.5 MB).

Appendix F (Continued):
Human Health Risk Assessment Supporting Documentation
Section: F.8
(pages 219-317)
is available
in a separate file (size: 3.8 MB).

Appendix F (Continued):
Human Health Risk Assessment Supporting Documentation
Section: F.9
(pages 318-403)
is available
in a separate file (size: 2.4 MB).

Appendix F (Continued):
Human Health Risk Assessment Supporting Documentation
Sections: F.10 through F.12
(pages 404-500)
are available
in a separate file (size: 2.8 MB).